

PATENT

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Appln. No. : 10/531,752
Applicant : Keith R. Minnich et al
International Appln. No. : PCT/US2003/033066
International Filing Date : October 17, 2003
Title : Method and Apparatus for High Efficiency Evaporation Operation
Docket No. : 4553-00013

) CERTIFICATE OF MAILING
) I hereby certify that this correspondence
) is being deposited with the United States
) Postal Service with sufficient postage as
) first class mail in an envelope addressed
) to: Commissioner for Patents, P.O. Box
) 1450, Alexandria, VA 22313-1450, on
) this 25th day of April, 2007.
)
)
) *Aleshia Prange 4-25-07*
) Aleshia Prange Date
)
)

RENEWED PETITION UNDER 37 C.F.R. §1.47(a)

Mail Stop: PCT
Commissioner for Patents
Office of PCT Legal Administration
P.O. Box 1450
Alexandria, VA 22313-1450

ATTN: Anthony Smith, Attorney Advisor
Office of PCT Legal Administration

Sir:

This is a request for reconsideration of the "Petition Under Rule 37 C.F.R. §1.47(a) to Accept Filing when Inventor Refuses to Sign" as filed June 12, 2006, and the Decision to dismiss that Petition mailed November 3, 2006. This paper is accompanied by a request for a four (4) month extension of time and requisite fees in a separate attachment appended hereto.

In the Decision dated November 3, 2006, it is stated, where the Office is being asked to accept the silence of the non-signing inventor as evidence of refusal to sign, Petitioner is required to provide some evidence that the application materials have been received by the non-signing applicant. Accordingly, Petitioner submits herewith a statement of facts from a person having firsthand knowledge of the facts that a complete

Application No. 10/531,752

Applicant: Keith R. Minnich et al

Renewed Petition Under Rule 37 C.F.R. §1.47(a) Dated April 25, 2007

copy of the application papers was sent to Ramkumar Karlapudi, and the date such papers were sent, along with a cover letter of instruction and proof of delivery evidence from DHL and Federal Express. Attached is a Statement of Facts in Support of Filing on Behalf of the Non-Signing Inventor as signed by Lisa Schollaert.

It is submitted that this Statement of Facts provides the necessary evidence from one having firsthand knowledge of the facts that the application materials including the Declaration were received by the non-signing inventor, and that his silence and lack of cooperation despite repeated attempts to contact him is evidence of his refusal to sign the Declaration.

Based on the foregoing, Petitioner requests that the Petition of June 12, 2006, be granted, so that a filing receipt can be obtained and the application can be advanced to examination.

Respectfully submitted,

ANDRUS, SCEALES, STARKE & SAWALL, LLP

By


William L. Falk
Reg. No. 27,709

Andrus, Sceales, Starke & Sawall, LLP
100 East Wisconsin Avenue, Suite 1100
Milwaukee, Wisconsin 53202
Telephone: (414) 271-7590
Facsimile: (414) 271-5770

PATENT

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Appln. No. : 10/531,752) CERTIFICATE OF MAILING
Applicant : Keith R. Minnich et al) I hereby certify that this correspondence
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Filing Date : October 17, 2003) to: Commissioner for Patents, P.O. Box
Title : Method and Apparatus for) 1450, Alexandria, VA 22313-1450, on
High Efficiency) this 25th day of April, 2007.
Evaporation Operation)
Docket No. : 4553-00013)
Aleshia Prange 4-25-07
Aleshia Prange Date

STATEMENT OF FACTS IN SUPPORT OF FILING ON BEHALF
OF NON-SIGNING INVENTOR UNDER 37 C.F.R. §1.47

Mail Stop: PCT
Commissioner for Patents
Office of PCT Legal Administration
P.O. Box 1450
Alexandria, VA 22313-1450

ATTN: Anthony Smith, Attorney Advisor
Office of PCT Legal Administration

Sir:

This statement is made as to the facts that are relied upon to establish the diligent effort made to secure the execution of the Declaration by the non-signing inventor, Mr. Ramkumar Karlapudi, for the above-identified patent application subsequent to filing thereof in the U.S. Patent and Trademark Office.

This statement is being made by the available person having firsthand knowledge of the facts recited herein. The person making this statement is:

Ms. Lisa Schollaert
Executive Assistant to the CEO
Aquatech International Corporation
1120 James Drive, Suite B-105
Hartland, Wisconsin 53029

Application No. 10/531,752

Applicant: Keith R. Minnich et al

Statement of Facts in Support of Filing on Behalf

of Non-Signing Inventor Under 37 C.F.R. §1.47 Dated 23 April - 2007

1. In an attempt to establish the latest last known location of the non-signing inventor, Mr. Karlapudi, I learned of his current work contact information from Mr. Devesh Mittal, an employee of Aquatech International Corporation on December 27, 2006. Through several e-mails directly with the non-signing inventor, I was able to determine the last known residential address of Mr. Karlapudi as: 1705 Cherry Ridge Drive, Lake Mary, Florida 32746.

Attached is Exhibit A, which includes copies of various e-mails showing the process of obtaining Mr. Karlapudi's last known home address.

2. On January 16, 2007, I sent via DHL to Ramkumar Karlapudi a cover letter of instruction along with a complete copy of the application including the specification, claims, drawings and Declaration as signed by the other inventors, Keith R. Minnich and Richard M. Schoen. I also sent a new Declaration to be signed by Mr. Karlapudi.

Attached is Exhibit B, which includes a copy of the cover letter, the complete copy of the application, the Declaration previously signed by the co-inventors, and the new Declaration.

Attached is Exhibit C, which includes DHL tracking delivery signature details and verifies that the enclosures of Exhibit B were delivered January 16, 2007, to Mr. Ramkumar Karlapudi at 11:13 a.m. and left at the front door at 1705 Cherry Ridge Drive, Lake Mary, Florida 32746.

3. On January 31 and March 13, 2007, I sent e-mails to Mr. Karlapudi asking if he would be signing the Declaration sent to him. These e-mails were not answered.

Attached is Exhibit D, which shows copies of these e-mails sent to Mr. Karlapudi.

4. On March 14, 2007, I sent via Federal Express to Ramkumar Karlapudi another copy of the cover letter and the application, the Declaration as signed by the previous inventors and a new Declaration (Exhibit B).

Attached is Exhibit E, which includes Federal Express tracking details and verifies that the enclosures of Exhibit B were delivered at 12:48 p.m. on March 15, 2007, and

Application No. 10/531,752

Applicant: Keith R. Minnich et al

Statement of Facts in Support of Filing on Behalf

of Non-Signing Inventor Under 37 C.F.R. §1.47 Dated 23 April 2007

received and signed for by K. Karlapudi at 1705 Cherry Ridge Drive, Lake Mary, Florida 32746.

No response has been received from Mr. Karlapudi despite two delivery attempts and various e-mails.

The above facts clearly demonstrate that despite being presented with the application and the inventor Declaration, Mr. Karlapudi refused to sign the inventor Declaration. Accordingly, the above facts demonstrate a bona fide and diligent effort was made to obtain the signature of Mr. Karlapudi on the inventor Declaration, but to no avail.

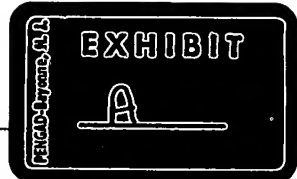
Dated: April 23, 2007



Lisa Schollaert

ATTORNEYS FOR APPLICANT:

Andrus, Sceales, Starke & Sawall, LLP
100 East Wisconsin Avenue, Suite 1100
Milwaukee, Wisconsin 53202
Telephone: (414) 271-7590
Facsimile: (414) 271-5770

Bill Falk

From: Lisa Schollaert [SchollaertL@aquatech.com]
Sent: Wednesday, December 27, 2006 3:51 PM
To: Bill Falk
Subject: RE: Location of Ram Kumar

Bill
 I will try first thing tomorrow
 Lisa

-----Original Message-----

From: Bill Falk [mailto:billf@andruslaw.com]
Sent: Wednesday, December 27, 2006 4:51 PM
To: Lisa Schollaert
Subject: RE: Location of Ram Kumar

Hi Lisa Thanks for the quick results!

Can you call Mr. Karlapudi, and refresh his memory about your past conversation earlier this year in which he appeared to offer his cooperation in signing the patent application. Ask him if you might e-mail a cover letter request along with a copy of the patent application enclosures sent November 28, 2005 by DHL, namely, the specification, claims, drawings and the Declaration(which he needs to sign and date). It would be great if he could return the fully signed Declaration as soon as possible.

Please let me know if you are able to do this.

Regards,
 Bill

From: Lisa Schollaert [mailto:SchollaertL@aquatech.com]
Sent: Wednesday, December 27, 2006 3:33 PM
To: Bill Falk
Subject: FW: Location of Ram Kumar

Found him!

-----Original Message-----

From: Devesh Mittal [mailto:mittald@aquatech.com]
Sent: Wednesday, December 27, 2006 3:47 PM
To: Lisa Schollaert
Cc: C. K. Tiwari
Subject: RE: Location of Ram Kumar

Lisa,

See below.

*Full Name: Ram Kumar Karlapudi
 Job Title: Director, Industrial Water
 Company: Pall Advanced Separations Systems*

Business Address: 1750 Filter Drive,

DeLand, FL 32724-2045

Business: (386) 822-8093
Business 2: (386) 822-8000 x393
Business Fax: (386) 822-8011

E-mail: rkarlapudi@pall.com
Web Page: <http://www.pall.com>

Best Regards,
Devesh

From: Lisa Schollaert [mailto:SchollaertL@aquatech.com]
Sent: Wednesday, December 27, 2006 12:52 PM
To: Devesh Mittal
Subject: FW: Location of Ram Kumar

Devesh
Can you help locate Ram Kumar for me?
Thanks

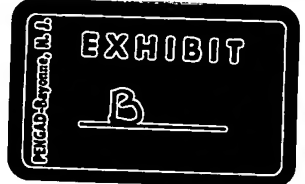
Lisa

-----Original Message-----

From: C. K. Tiwari [mailto:tiwaric@aquatech.com]
Sent: Wednesday, December 27, 2006 1:45 PM
To: Lisa Schollaert
Cc: Venkee Sharma
Subject: RE: Location of Ram Kumar

I had seen a mail from Devesh Mittal stating that he ran into Ram during Powergen in Florida. It is my understanding that he has moved to FL and Devesh may be able to tell you whether he exchanged any business cards with him or not.

Best Regards
C. K. Tiwari
Aqua-chem ICD Division
Aquatech International Corporation
1120 James Dr., Suite B 105
Hartland, WI 53029
P) 262-369-9595 ext 120
F) 262-369-9494
e-mail - tiwaric@aquatech.com
From innovation flows leadership



Aquatech
International Corporation

One - Four Coins Drive
Canonsburg, PA 15317 USA
t: 724 746 5300
f: 724 746 5359
www.aquatech.com

Ramkumar Karlapudi
1705 Cherry Ridge Drive
Lake Mary, FL 32746

Re: U.S. Patent Application Serial No. 10/531,752
Our Ref.: 4553-00013

Dear Ramkumar:

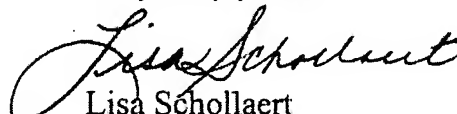
Enclosed for your review is a full copy of U.S. Patent Application Serial No. 10/531,752 as filed April 18, 2005. The application includes the specification, claims, drawings and Declaration as signed by the other inventors Keith R. Minnich and Richard M. Schoen. Also enclosed is a copy of the claims as further revised in a Preliminary Amendment filed April 18, 2005.

Please sign and date the enclosed new Declaration where indicated and return the executed Declaration to me as soon as possible in the enclosed self-addressed, stamped envelope.

If you have any questions, please contact me.

I thank you in advance for your cooperation.

Very truly yours,


Lisa Schollaert
Executive Assistant to the CEO

Under the Paperwork Reduction Act of 1995, no persons are required to respond to a collection of information unless it displays a valid OMB control number.

TRANSMITTAL LETTER TO THE UNITED STATES DESIGNATED/ELECTED OFFICE (DO/EO/US) CONCERNING A SUBMISSION UNDER 35 U.S.C. 371		ATTORNEY'S DOCKET NUMBER 4553-00013
INTERNATIONAL APPLICATION NO. PCT/US2003/033066		INTERNATIONAL FILING DATE 17 October 2003
TITLE OF INVENTION Method and Apparatus for High Efficiency Evaporation Operation		
APPLICANT(S) FOR DO/EO/US		
<p>Applicant herewith submits to the United States Designated/Elected Office (DO/EO/US) the following items and other information:</p> <ol style="list-style-type: none"> 1. <input checked="" type="checkbox"/> This is a FIRST submission of items concerning a submission under 35 U.S.C. 371. 2. <input type="checkbox"/> This is a SECOND or SUBSEQUENT submission of items concerning a submission under 35 U.S.C. 371. 3. <input type="checkbox"/> This is an express request to begin national examination procedures (35 U.S.C. 371(f)). The submission must include items (5), (6), (9) and (21) indicated below. 4. <input checked="" type="checkbox"/> The US has been elected (Article 31). 5. <input checked="" type="checkbox"/> A copy of the International Application as filed (35 U.S.C. 371(c)(2)) <ol style="list-style-type: none"> a. <input checked="" type="checkbox"/> is attached hereto (required only if not communicated by the International Bureau). b. <input checked="" type="checkbox"/> has been communicated by the International Bureau. c. <input type="checkbox"/> is not required, as the application was filed in the United States Receiving Office (RO/US). 6. <input type="checkbox"/> An English language translation of the International Application as filed (35 U.S.C. 371(c)(2)). <ol style="list-style-type: none"> a. <input type="checkbox"/> is attached hereto. b. <input type="checkbox"/> has been previously submitted under 35 U.S.C. 154(d)(4). 7. <input checked="" type="checkbox"/> Amendments to the claims of the International Application under PCT Article 19 (35 U.S.C. 371(c)(3)) <ol style="list-style-type: none"> a. <input type="checkbox"/> are attached hereto (required only if not communicated by the International Bureau). b. <input type="checkbox"/> have been communicated by the International Bureau. c. <input type="checkbox"/> have not been made; however, the time limit for making such amendments has NOT expired. d. <input checked="" type="checkbox"/> have not been made and will not be made. 8. <input type="checkbox"/> An English language translation of the amendments to the claims under PCT Article 19 (35 U.S.C. 371(c)(3)). 9. <input type="checkbox"/> An oath or declaration of the inventor(s) (35 U.S.C. 371(c)(4)). 10. <input type="checkbox"/> An English language translation of the annexes of the International Preliminary Examination Report under PCT Article 36 (35 U.S.C. 371(c)(5)). <p>Items 11 to 20 below concern document(s) or information included:</p> <ol style="list-style-type: none"> 11. <input type="checkbox"/> An Information Disclosure Statement under 37 CFR 1.97 and 1.98. 12. <input type="checkbox"/> An assignment document for recording. A separate cover sheet in compliance with 37 CFR 3.28 and 3.31 is included. 13. <input checked="" type="checkbox"/> A preliminary amendment. 14. <input type="checkbox"/> An Application Data Sheet under 37 CFR 1.76. 15. <input type="checkbox"/> A substitute specification. 16. <input type="checkbox"/> A power of attorney and/or change of address letter. 17. <input type="checkbox"/> A computer-readable form of the sequence listing in accordance with PCT Rule 13ter.2 and 37 CFR 1.821- 1.825. 18. <input type="checkbox"/> A second copy of the published International Application under 35 U.S.C. 154(d)(4). 19. <input type="checkbox"/> A second copy of the English language translation of the international application under 35 U.S.C. 154(d)(4). 20. <input checked="" type="checkbox"/> Other items or information: Supplement to Transmittal Letter; International Search Report 		

This collection of information is required by 37 CFR 1.414 and 1.491-1.492. The information is required to obtain or retain a benefit by the public, which is to file (and by the USPTO to process) an application. Confidentiality is governed by 35 U.S.C. 122 and 37 CFR 1.11 and 1.14. This collection is estimated to take 15 minutes to complete, including gathering information, preparing, and submitting the completed form to the USPTO. Time will vary depending upon the individual case. Any comments on the amount of time you require to complete this form and/or suggestions for reducing this burden, should be sent to the Chief Information Officer, U.S. Patent and Trademark Office, U.S. Department of Commerce, P.O. Box 1450, Alexandria, VA 22313-1450. DO NOT SEND FEES OR COMPLETED FORMS TO THIS ADDRESS. SEND TO: Mail Stop PCT, Commissioner for Patents, P.O. Box 1450, Alexandria, VA 22313-1450.

Under the Paperwork Reduction Act of 1995, no persons are required to respond to a collection of information unless it displays a valid OMB control number.

U.S. APPLICATION NO. (if known, see 37 CFR 1.5)		INTERNATIONAL APPLICATION NO. PCT/US2003/033066		ATTORNEY'S DOCKET NUMBER 4553-00013																																																																																																																																					
<p>The following fees have been submitted</p> <table border="1"> <tr> <td>21. <input checked="" type="checkbox"/> Basic national fee.....</td> <td>\$300</td> <td colspan="2">CALCULATIONS</td> <td colspan="2">PTO USE ONLY</td> </tr> <tr> <td>22. <input checked="" type="checkbox"/> Examination fee If International preliminary examination report prepared by USPTO and all claims satisfy provisions of PCT Article 33(1)-(4).....</td> <td>\$100</td> <td>\$</td> <td>300.00</td> <td></td> <td></td> </tr> <tr> <td>All other situations.....</td> <td>\$200</td> <td>\$</td> <td>200.00</td> <td></td> <td></td> </tr> <tr> <td>23. <input checked="" type="checkbox"/> Search fee Search fee (37 CFR 1.445(a)(2)) has been paid on the international application to the USPTO as an International Searching Authority.....</td> <td>\$100</td> <td>\$</td> <td>400.00</td> <td></td> <td></td> </tr> <tr> <td>International Search Report prepared and provided to the Office.....</td> <td>\$400</td> <td>\$</td> <td>900.00</td> <td></td> <td></td> </tr> <tr> <td>All other situations.....</td> <td>\$500</td> <td></td> <td></td> <td></td> <td></td> </tr> </table> <p>TOTAL OF 21, 22 and 23 =</p> <table border="1"> <tr> <td><input type="checkbox"/> Additional fee for specification and drawings filed in paper over 100 sheets (excluding sequence listing or computer program listing filed in an electronic medium). The fee is \$250 for each additional 50 sheets of paper or fraction thereof.</td> <td></td> <td></td> <td></td> <td></td> <td></td> </tr> <tr> <td>Total Sheets</td> <td>Extra Sheets</td> <td>Number of each additional 50 or fraction thereof (round up to a whole number)</td> <td>RATE</td> <td></td> <td></td> </tr> <tr> <td>50 - 100 = 0</td> <td>/50 = 0</td> <td></td> <td>x \$250.00</td> <td>\$ 0.00</td> <td></td> </tr> </table> <p>Surcharge of \$130.00 for furnishing the oath or declaration later than 30 months from the earliest claimed priority date (37 CFR 1.492(h)).</p> <table border="1"> <tr> <td>CLAIMS</td> <td>NUMBER FILED</td> <td>NUMBER EXTRA</td> <td>RATE</td> <td>\$ 900.00</td> <td></td> </tr> <tr> <td>Total claims</td> <td>137 - 20 =</td> <td>107</td> <td>x \$50.00</td> <td>\$ 5,350.00</td> <td></td> </tr> <tr> <td>Independent claims</td> <td>5 - 3 =</td> <td>2</td> <td>x \$200.00</td> <td>\$ 400.00</td> <td></td> </tr> <tr> <td>MULTIPLE DEPENDENT CLAIM(S) (if applicable)</td> <td></td> <td></td> <td>+ \$360.00</td> <td>\$ 360.00</td> <td></td> </tr> <tr> <td colspan="4">TOTAL OF ABOVE CALCULATIONS =</td> <td>\$ 7,010.00</td> <td></td> </tr> <tr> <td colspan="4"><input checked="" type="checkbox"/> Applicant claims small entity status. See 37 CFR 1.27. Fees above are reduced by ½.</td> <td>3,505.00</td> <td></td> </tr> <tr> <td colspan="4"></td> <td>SUBTOTAL =</td> <td>\$ 3,505.00</td> </tr> <tr> <td colspan="4">Processing fee of \$130.00 for furnishing the English translation later than 30 months from the earliest claimed priority date (37 CFR 1.492(i)).</td> <td>\$ 0.00</td> <td></td> </tr> <tr> <td colspan="4"></td> <td>TOTAL NATIONAL FEE =</td> <td>\$ 3,505.00</td> </tr> <tr> <td colspan="4">Fee for recording the enclosed assignment (37 CFR 1.21(h)). The assignment must be accompanied by an appropriate cover sheet (37 CFR 3.28, 3.31). \$40.00 per property</td> <td>\$ 0.00</td> <td></td> </tr> <tr> <td colspan="4"></td> <td>TOTAL FEES ENCLOSED =</td> <td>\$ 3,505.00</td> </tr> <tr> <td colspan="4"></td> <td>Amount to be refunded:</td> <td>\$</td> </tr> <tr> <td colspan="4"></td> <td>Amount to be charged:</td> <td>\$</td> </tr> </table> <p>a. <input checked="" type="checkbox"/> A check in the amount of \$ 3,505.00 to cover the above fees is enclosed.</p> <p>b. <input type="checkbox"/> Please charge my Deposit Account No. 01.2000 in the amount of \$ _____ to cover the above fees. A duplicate copy of this sheet is enclosed.</p> <p>c. <input checked="" type="checkbox"/> The Commissioner is hereby authorized to charge any additional fees which may be required, or credit any overpayment to Deposit Account No. 01.2000. A duplicate copy of this sheet is enclosed.</p> <p>d. <input type="checkbox"/> Fees are to be charged to a credit card. WARNING: Information on this form may become public. Credit card information should not be included on this form. Provide credit card information and authorization on PTO-2038.</p> <p>NOTE: Where an appropriate time limit under 37 CFR 1.495 has not been met, a petition to revive (37 CFR 1.137(a) or (b)) must be filed and granted to restore the International Application to pending status.</p> <p>SEND ALL CORRESPONDENCE TO: Andrus, Sceales, Stark & Sawall, LLP, 100 East Wisconsin Avenue, Suite 1100, Milwaukee, WI 53202-4178, (414) 271-7590, Facsimile: (414) 271-5770</p> <p>CERTIFICATE OF EXPRESS MAIL: I hereby certify that this correspondence is being deposited with the United States Postal Service with sufficient postage as EXPRESS MAIL-POST OFFICE TO ADDRESSEE, in an envelope addressed to: BOX PCT COMMISSIONER OF PATENTS, P.O. Box 1450, Alexandria, VA 22313-1450 on 18 April 2005</p> <p>The Express Mail Label is EV477160526US</p> <p><i>Thomas M. Wozny</i> SIGNATURE Thomas M. Wozny NAME 28,922 REGISTRATION NUMBER</p> <p><i>Dorothy A. Hauser</i> April 18, 2005 Dorothy A. Hauser Date</p>						21. <input checked="" type="checkbox"/> Basic national fee.....	\$300	CALCULATIONS		PTO USE ONLY		22. <input checked="" type="checkbox"/> Examination fee If International preliminary examination report prepared by USPTO and all claims satisfy provisions of PCT Article 33(1)-(4).....	\$100	\$	300.00			All other situations.....	\$200	\$	200.00			23. <input checked="" type="checkbox"/> Search fee Search fee (37 CFR 1.445(a)(2)) has been paid on the international application to the USPTO as an International Searching Authority.....	\$100	\$	400.00			International Search Report prepared and provided to the Office.....	\$400	\$	900.00			All other situations.....	\$500					<input type="checkbox"/> Additional fee for specification and drawings filed in paper over 100 sheets (excluding sequence listing or computer program listing filed in an electronic medium). 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IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re the Application of:

Keith R. Minnich

Ramkumar Karlapudi

Richard M. Schoen

Filed Herein

International Appl. No.

PCT/US2003/033066

International Filing Date:

17 October 2003

Priority Date Claimed:

18 October 2002

Method and Apparatus for High
Efficiency Evaporation Operation

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Dorothy A. Hauser

April 18, 2005

Date

SUPPLEMENT TO TRANSMITTAL LETTER

Mail Stop: PCT
Commissioner for Patents
P.O. Box 1450
Alexandria, VA 22313-1450

Sir:

This application is being filed to enter the national stage of the PCT pursuant to 37 C.F.R. §1.495(c). The application includes the specification, at least one claim, and the filing fee. The Declaration is not included. The inventors are

- 1) Keith R. Minnich
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- 2) Ramkumar Karlapudi
1981 Foxcroft Lane
Waukesha, WI 53189
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- 3) Richard M. Schoen
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Hartland, WI 53029
US citizen

Applicant: Keith R. Minnich et al
Attorney Docket No.: 4553-00013

The Declaration will be forwarded promptly upon notification from the U.S. Patent and Trademark Office.

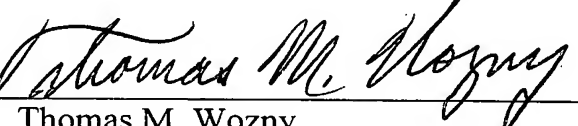
Please address all correspondence and telephone calls regarding this matter to:

Thomas M. Wozny
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(414) 271-7590

Respectfully submitted,

ANDRUS, SCEALES, STARKE & SAWALL, LLP

By


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IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re the Application of:

Keith R. Minnich

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Dorothy A. Hauser April 18, 2005

Date

Mail Stop: PCT
Commissioner for Patents
P.O. Box 1450
Alexandria, VA 22313-1450

Sir:

In the matter of the above-identified patent application, please enter the following:

Amendments to the Specification begin on page 2 of this paper.

Amendments to the Claims are reflected in the listing of claims which begins on page 3 of this paper.

There are no Amendments to the Drawings made in this paper.

Remarks begin on page 18 of this paper.

Applicant: Keith R. Minnich et al
Attorney Docket No.: 4553-00013

In the Specification:

Please add the following heading and paragraph at page 1, between the title and the first line of text as follows:

CROSS-REFERENCE TO RELATED APPLICATIONS

This application is the U.S. national stage application of International Application PCT/US2003/03306, filed 17 October 2003, which international application was published on 29 April 2004, as International Publication WO2004/035479 in the English language. The International Application claims priority of United States Provisional Application No 60/419,549 filed on 18 October 2002, and United States Provisional Application No 60/419,552 filed on 18 October 2002.

Please add the enclosed "Abstract of the Disclosure" as new page 44 to the specification.

Applicant: Keith R. Minnich et al
Attorney Docket No.: 4553-00013

In the Claims:

Please rewrite claim 4, 40, 44, 54 and 97 as follows:

1. (Original) A process for treatment of an aqueous feed stream in heat transfer equipment, said heat transfer equipment comprising at least one evaporator with at least one heat transfer surface, to produce a low solute containing distillate stream and a high solute/solids containing blowdown stream, in which said feed stream has minimal tendency to scale said heat transfer surface, said process comprising:

(a) providing a feed water stream containing soluble and insoluble inorganic and organic species therein, said species comprising:

(II) multi-valent metal cations,

(III) alkalinity,

(IV) at least one molecular species which is at low ionization levels when in solution at around neutral pH;

(b) removing a portion or substantially all multi-valent cations metal from said feed stream, and

(c) reducing the tendency of said feed water or form scale on said heat transfer surfaces, when said feed water is concentrated to a selected concentration factor at a selected pH, by effecting, in any order, one or more of the following;

(I) removing substantially all alkalinity from said feed water stream;

(II) removing dissolved gas from said feed water stream;

(III) raising the pH of said feed water stream to at least about 9 or higher;

(d) passing the product from step (c) into said heat transfer equipment, wherein said heat transfer equipment:

(I) contains a plurality of heat transfer surfaces,

(II) contains a circulating high solids solution, and

(III) wherein the pH of said circulating solution is maintained to at least about 9, or higher,

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(e) so as to concentrate said feed water to said selected concentration factor, to produce:

- (I) a high solute/solids containing blowdown stream, and
- (II) a low solute containing distillate stream.

2. (Original) The process as set forth in claim 1, wherein the step of removing said multi-valent cations includes removing substantially all said alkalinity associated with hardness, and is accomplished in a single unit operation.

3. (Original) The process as set forth in claim 2, wherein said single unit operation comprises a weak acid cation ion exchange system operated in a hydrogen form.

4. (Currently Amended) The process as set forth in claim 4~~(b)~~1, wherein the step (b) of said multi-valent cation removal is accomplished in a weak acid cation ion exchange system that is operated in a sodium form.

5. (Original) The process as set forth in claim 1, further comprising the step of adding acid before the step of removing said dissolved gas, to effect conversion of alkalinity to carbon dioxide.

6. (Original) The process as set forth in claim 3, wherein said feed water stream contains more multi-valent cations than alkalinity, and further comprising, before feeding said feed water to said weak acid cation exchange system, the step of adjusting the ratio of multi-valent cations to alkalinity by adding a base to said feed water, so as to raise the alkalinity of said feed water.

7. (Original) The process as set forth in claim 3, wherein said feed water stream contains more alkalinity than multi-valent cations, and further comprising, before feeding said feed water to said weak acid cation exchange system, the step of addition of acid to said feed water, so as to remove the excess alkalinity in said feed water.

8. (Original) The process as set forth in claim 1, wherein the step of multi-valent cation removal is accomplished by passing said feed water through a sodium form strong acid cation ion exchange system.

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9. (Original) The process as set forth in claim 1, wherein in step (c), the pH is raised to between 10 and 11.

10. (Original) The process as set forth in claim 1, wherein in step (c), the pH is raised to between 9 and 10.

11. (Original) The process as set forth in claim 10, wherein said sparingly ionized species when in neutral or near neutral pH aqueous solution comprises a weak acid with a pK_{a_1} of about 7.5 or higher.

12. (Original) The process as set forth in claim 1, wherein in step (c), the pH is raised to between 11 and 12.

13. (Original) The process as set forth in claim 1, wherein in step (c), the pH is raised to between 12 and 13.

14. (Original) The process as set forth in claim 1, wherein in step (c), the pH is raised to greater than or at least about 13.

15. (Original) The process as set forth in claim 1, wherein said low ionized species when in neutral or near neutral pH aqueous feed stream comprises silica (SiO_2).

16. (Original) The process as set forth in claim 1, wherein said low ionized species when in neutral or near neutral pH aqueous feed stream comprises meta/ortho silicic acid ($H_4 SiO_4$).

17. (Original) The process as set forth in claim 1, wherein said low ionized species when in neutral or near neutral pH aqueous feed stream comprises an ionizable organic carbon species.

18. (Original) The process as set forth in claim 1, wherein said low ionized species when in neutral or near neutral pH aqueous feed stream comprises boron, or derivatives thereof.

19. (Original) The process according to claim 1, wherein the step of removal of multi-valent cations is accomplished by addition of an alkali to simultaneously raise pH while precipitating hardness from said feed water stream.

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20. (Original) The process as set forth in claim 1, further comprising the step of adding caustic before the step of removing dissolved gas, to effect removal of gasses such as ammonia.

21. (Original) The process as set forth in claim 1, wherein said at least one molecular species which is at low ionization levels when in solution at around neutral pH comprises silica, and wherein said blowdown stream contains silica up to about 160 ppm.

22. (Original) The process as set forth in claim 21, wherein said silica is present in said blowdown stream at up to about 2500 ppm.

23. (Original) The process as set forth in claim 21, wherein said silica is present in said blowdown stream at up to about 5000 ppm.

24. (Original) The process as set forth in claim 21, wherein said silica is present in said blowdown stream at up to about 7500 ppm.

25. (Original) The process as set forth in claim 21, wherein said silica is present in said blowdown stream at up to about 10,000 ppm.

26. (Original) The process as set forth in claim 21, wherein said silica is present in said blowdown stream at up to about 20,000 ppm.

27. (Original) The process as set forth in claim 21, wherein said silica is present in said blowdown stream at up to about 50,000 ppm.

28. (Original) The process as set forth in claim 21, wherein said silica is present in said blowdown stream at up to about 75,000 ppm.

29. (Original) The process as set forth in claim 21, wherein said silica is present in said blowdown stream at up to about 100,000 ppm.

30. (Original) The process as set forth in claim 1, wherein said feed water stream comprises silica, and wherein said heat transfer equipment is operated without limitation of the concentration of silica present in said blowdown stream.

31. (Original) The process according to claim 1, wherein the ratio of the quantity of said distillate stream produced to the quantity of said feed water stream provided is greater than about 50%.

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32. (Original) The process according to claim 1, wherein the ratio of the quantity of said distillate stream produced to the quantity of said feed water stream provided is greater than about 85%.

33. (Original) The process according to claim 1, wherein the ratio of the quantity of said distillate stream produced to the quantity of said feed water stream provided is greater than about 95%.

34. (Original) The process according to claim 1, wherein the ratio of the quantity of said distillate stream produced to the quantity of said feed water stream provided is greater than about 99%.

35. (Original) The process according to claim 1, wherein said heat transfer equipment comprises falling thin film evaporation equipment, operating as a single unit, or operating in series, or operating in parallel to generate said distillate stream and said blowdown stream.

36. (Original) The process according to claim 1, wherein said heat transfer equipment comprises forced circulation evaporation equipment operating as a single unit or operating in parallel to generate said distillate stream and a high solids blowdown stream.

37. (Original) The process according to claim 1, wherein said heat transfer equipment comprises natural circulation evaporation equipment operating as a single unit or operating in parallel to generate said distillate stream and a high solids blowdown stream.

38. (Original) The process according to claim 35 or claim 36 or claim 37, wherein said heat transfer surfaces are tubular.

39. (Original) The process according to claim 35 or claim 36 or claim 37, wherein said heat transfer surfaces are plates.

40. (Currently Amended) The process as set forth in claim 38 or claim 39, wherein said heat transfer surfaces are operated in a vertical position.

41. (Original) The process as set forth in claim 38, wherein said heat transfer surfaces are operated in a horizontal position.

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42. (Original) The process as set forth in claim 38 wherein said heat transfer surfaces are designed for enhanced heat transfer.

43. (Original) The process as set forth in claim 38 wherein said circulating solution is heated on the interior of the tubes.

44. (Currently Amended) The process as set forth in claim 38 or claim 39, wherein said circulating solution is heated on the exterior of the tubes or plates.

45. (Original) The process as set forth in claim 39, wherein said plates are die pressed plates.

46. (Original) The process as set forth in claim 39, wherein said plates are made from flat sheets welded together and then formed into final shape by internal pressure.

47. (Original) The process as set forth in claim 39, wherein said plates are fabricated as welded assemblies.

48. (Original) The process as set forth in claim 39, wherein said plates are gasketed.

49. (Original) The process as set forth in claim 35 or claim 36 or claim 37, wherein said heat transfer equipment is operated in a steam driven multiple effect mode.

50. (Original) The process as set forth in claim 35 or claim 36 or claim 37, wherein said heat transfer equipment is operated in a mechanical vapor recompression mode.

51. (Original) The process as set forth in claim 35 or claim 36 or claim 37, wherein said heat transfer equipment is operated in a thermal compression mode.

52. (Original) The process as set forth in claim 35 or claim 36 or claim 37, wherein said heat transfer equipment is operated as a multiple stage flash evaporator.

53. (Original) The process as set forth in claim 35 or claim 36 or claim 37, further comprising the step of treating said high solute concentrate stream in a crystallizer operating as a single unit or operating in parallel to generate said distillate stream and said high solids blowdown stream.

54. (Currently Amended) The process as set forth in claim 36 or ~~claim 53~~, comprising the step of further treating said high solids blowdown stream after reaching said selected concentration factor in a solids dewatering device operating as a single unit or operating in parallel.

55. (Original) The process as set forth in claim 54, further comprising the step of generation of a high solute low suspended solids filtrate water stream, and still further comprising the step of directing said low filtrate water stream to the inlet of said heat transfer equipment for further processing.

56. (Original) The process as set forth in claim 55, further comprising the step of generation of a dry solids product for disposal.

57. (Original) The process as set forth in claim 1, wherein the step of removing said dissolved gases, is further comprised of lowering the pH of said feed stream to remove any remaining alkalinity and release carbon dioxide.

58. (Original) The process as set forth in claim 57, further comprised of heating said acidified feed water stream to enhance gas removal in a degasifier prior to entering said heat transfer equipment.

59. (Original) The process as set forth in claim 1, wherein step (d) further comprising distributing said circulating solution across one side of said plurality of heat transfer surfaces to generate a steam vapor.

60. (Original) The process as set forth in claim 59, further comprising collecting said steam vapor and slightly compressing it to form a compressed steam vapor.

61. (Original) The process as set forth in claim 60, further comprising directing said compressed steam vapor to a second side of said plurality of heat transfer surfaces to condense said compressed steam vapor into said distillate stream.

62. (Original) The process according to claim 1, wherein the step of raising the pH is accomplished by addition of a base in aqueous solution, said base selected from the group consisting of (a) sodium hydroxide, (b) sodium carbonate, (c) potassium hydroxide, and (d) potassium carbonate.

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(III) minimizing gases dissolved or suspended in said aqueous feed stream;

(b) then, after step (a), increasing the pH of said aqueous feed stream in said evaporation equipment to at least about 9, or higher.

92. (Original) The process as set forth in claim 1, or claim 78, or claim 90, or claim 91, further comprising, during the step of removing alkalinity, the additional step of removing substantially all non-hydroxide alkalinity not associated with hardness.

93. (Original) The process as set forth in claim 90, or claim 91, wherein the step of raising the pH of said feed water stream, comprises raising the pH to between about 10 and about 11.

94. (Original) The process as set forth in claim 90, or claim 91, wherein the step of raising the pH of said feed water stream comprises raising the pH to between about 11 and about 12.

95. (Original) The process as set forth in claim 90, or claim 91, wherein the step of raising the pH of said feed water stream comprises raising the pH to between about 12 and about 13.

96. (Original) The process according to claim 1, wherein said feed water stream comprises effluents from hydrocarbon recovery operations as produced water.

97. (Currently Amended) The process as set forth in claim 90, or claim 91, or claim 91, wherein during the step of raising the pH of said feed water stream comprises raising the pH to between about 9 and about 10.

98. (Original) The process as set forth in claim 19, wherein a high solids containing waste stream is generated and, further comprising, de-watering of said high solids containing waste stream.

99. (Original) The process as set forth in claim 98, wherein a low suspended solids stream is generated and, further comprising directing said low suspended solids stream back to the inlet of said softener.

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63. (Original) The process according to claim 1, wherein the step of raising the pH is accomplished by addition of an aqueous organic base.

64. (Original) The process according to claim 1, wherein the ratio of the quantity of said distillate stream produced to the quantity of said feed water stream provided is between 90 and 98 percent.

65. (Original) The process according to claim 1, wherein said feed water stream further comprises cooling tower blowdown.

66. (Original) The process according to claim 1, wherein said feed water further comprises scrubber blowdown.

67. (Original) The process according to claim 1, wherein said feed water further comprises water utilized in ash transport in a coal fired steam-electric power plant.

68. (Original) The process according to claim 1, wherein said feed water stream comprises ash pond water.

69. (Original) The process according to claim 1, wherein said feed water stream comprises ash-sluicing water.

70. (Original) The process according to claim 1, wherein said feed water stream comprises effluent from sewage treatment.

71. (Original) The process according to claim 1, wherein said feed water stream comprises effluent from a food processing treatment.

72. (Original) The process according to claim 1, wherein said feed water stream comprises boiler blowdown.

73. (Original) The process according to claim 1, wherein said feed water stream comprises a concentrated stream from membrane separation equipment.

74. (Original) The process according to claim 1, wherein said feed water stream comprises effluent from oil refining operations.

75. (Original) The process as set forth in claim 35 or claim 36 or claim 37, comprising the step of further treating said high solids blowdown stream after reaching said selected concentration factor in a spray dryer to dry solids.

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76. (Original) The process as set forth in claim 1, wherein the step of multi-valent cation removal is partially accomplished by passing said feed water stream through membrane softening equipment.

77. (Original) The process as set forth in claim 1, wherein the step of multi-valent cation removal is accomplished by increasing the pH to at least about 10 in said feed water stream and passing the pH adjusted stream through membrane separation equipment to filter out hardness precipitate.

78. (Original) In a process for the concentration of an aqueous feed stream in an evaporator to produce a low solute containing distillate stream, and retaining at least a portion of said aqueous feed stream in said evaporator to increase the concentration of a solute to a selected concentration factor at a selected pH in said retained portion of said aqueous feed stream, the improvement which comprises feeding said evaporator with an aqueous feed stream characterized at the time of initial entry into said evaporator, by:

- (I) substantially no multi-valent cations,
- (II) substantially no alkalinity,
- (III) substantially no dissolved or suspended gases, and
- (IV) a pH of at least 9 or above.

79. (Original) The process as set forth in claim 78 wherein greater than 80% of the multi-valent cations are removed from said feed stream.

80. (Original) The process as set forth in claim 78 wherein greater than 90% of the multi-valent cations are removed from said feed stream.

81. (Original) The process as set forth in claim 78 wherein greater than 98% of the multi-valent cations are removed from said feed stream.

82. (Original) The process as set forth in claim 78 wherein greater than 80% of the alkalinity is removed from said feed stream.

83. (Original) The process as set forth in claim 78 wherein greater than 90% of the alkalinity is removed from said feed stream.

84. (Original) The process as set forth in claim 78 wherein greater than 98% of the alkalinity is removed from said feed stream.

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85. (Original) The process as set forth in claim 78 wherein greater than 80% of the gases are removed from said feed stream.

86. (Original) The process as set forth in claim 78 wherein greater than 90% of the gases are removed from said feed stream.

87. (Original) The process as set forth in claim 78 wherein greater than 98% of the gases are removed from said feed stream.

88. (Original) The process as set forth in claim 78 wherein said bases are selected from the group consisting of carbon dioxide, ammonia, oxygen, nitrogen and mixtures thereof.

89. (Original) The process as set forth in claim 78, wherein said aqueous feed stream is further characterized by minimizing or eliminating scale inhibitor solution or scale dispersant solution in said aqueous feed stream.

90. (Original) A method for treating a feed water stream, in a least one evaporator system and for simultaneously (a) reducing the scaling potential and (b) allowing an increased heat transfer rate and (c) allowing higher recovery and (d) minimizing or eliminating scale control methods in an existing heat transfer system, to produce a low solute containing distillate stream and a high solute/solids containing blowdown stream, said method comprising:

(a) providing a feed water stream containing soluble and insoluble species therein, said species comprising two or more of the following:

(I) multi-valent metal cations,

(II) alkalinity, and

(III) at least one molecular species which is at low ionization levels when in solution at around neutral pH;

(b) removing a portion or substantially all multi-valent cations from said feed stream, and

(c) reducing the tendency of said feed water to form scale when said feed water is concentrated to a selected concentration factor at a selected pH, by effecting, in any order, one or more of the following;

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- (I) removing substantially all alkalinity from said feed water stream;
 - (II) removing dissolved or suspended gases from said feed water stream, whether initially present or created during said multi-valent cation or said alkalinity removal step or said pH adjustment step;
 - (III) raising the pH of said feed water stream to at least about 9 or higher;
- (d) passing the product from step (c) into heat transfer equipment, wherein said heat transfer equipment:
- (I) contains a plurality of heat transfer surfaces,
 - (II) contains a circulating high solutes/solids solution, and
 - (III) the pH of said circulating solution is maintained to at least about 9, or higher,
- (e) so as to concentrate said feed water to said selected concentration factor, to produce:
- (I) a high solute/solids containing blowdown stream, and
 - (II) a low solute containing distillate stream.

91. (Original) In a process for the purification of an aqueous feed stream comprising solutes and solvent by using evaporation equipment to increase the concentration of said aqueous stream to a selected concentration factor by generating a low solute containing distillate stream and retaining at least a portion of said aqueous feed stream in said evaporation equipment to increase the concentration of a selected solute to a selected concentration factor in said retained portion of said aqueous feed stream, the improvement which comprises controlling solutes, multi-valent metal cations, alkalinity, and carbon dioxide in said aqueous feed stream to a level where the tendency to form scale is effectively eliminated at said selected concentration factor, by

- (a) prior to feeding of said aqueous feed stream to said evaporation equipment, in any order,
- (I) minimizing multi-valent cations in said aqueous feed stream,
 - (II) minimizing alkalinity of said aqueous feed stream,

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100. (Original) The process as set forth in claim 90, or claim 91, wherein the step of raising the pH of said feed water stream comprises raising the pH to greater than or at least about 13.

101. (Original) The process as set forth in claim 1, wherein the steps of (b) removing multi-valent cations, and (c) removing alkalinity, removing dissolved gases, and increasing pH are accomplished prior to a membrane process to pre-concentrate the feed stream upstream of said heat transfer equipment described under step (d).

102. (Original) The process as set forth in claim 1, wherein the removal of multi-valent cations and partially raising the pH are accomplished prior to pre-concentrating said feed stream in a membrane process prior to step (c).

103. (Original) Apparatus for treatment of a feed water stream, said feed water stream characterized by the presence of two or more of the following:

(I) multi-valent metal cations,

(II) alkalinity,

(III) at least one molecular species which is at low ionization levels when in solution at around neutral pH, to produce a low solute containing distillate stream and a high solute/solids containing blowdown stream, said apparatus comprising:

a) pretreatment equipment for effectively eliminating the tendency of said feed water to form scale on heat transfer surfaces when said feed water is concentrated to a desired concentration factor at a selected pH, comprising, in any order:

(I) at least one softener for removing a portion or substantially all multi-valent cations from said feed stream, and one or more of the following:

(II) at least one de-alkalizer for removing essentially all alkalinity from said feed water stream,

(III) a degasifier for removing dissolved gases,

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(IV) chemical addition apparatus for raising the pH of said circulating solution in said heat transfer equipment to a selected pH of at least about 9 by adding a selected base thereto, to urge said at least one molecular species with low ionization levels when in solution at about neutral pH toward increased ionization;

(b) one or more evaporator units, said one or more evaporator units, treating said feed water to produce a high solute/solids containing blowdown stream and a low solute containing distillate stream, and to concentrate said feed water to said selected concentration factor.

104. (Original) The apparatus as set forth in claim 103, further comprising, downstream of one or more said evaporator units, to further process said low solute containing distillate stream therefrom, a cation exchange unit.

105. (Original) The apparatus as set forth in claim 103, further comprising, downstream of one or more said evaporator units, to further process said low solute containing distillate stream therefrom, an anion exchange unit.

106. (Original) The apparatus as set forth in claim 103, further comprising, downstream of one or more said evaporator units, to further process said low solute containing distillate stream therefrom, at least one mixed bed ion exchange unit.

107. (Original) The apparatus as set forth in claim 104 or claim 105 or claim 106, further comprising an ion exchange resin regenerator that generates an ion exchange regenerant stream, and further comprising means for directing said ion exchange regenerant stream to the inlet of said degasifier unit in order to treat said ion exchange regenerant stream in said evaporator.

108. (Original) The apparatus as set forth in claim 103, further comprising, downstream of one or more said evaporator units, to further process the said low solute containing distillate stream therefrom, a continuous electrodeionization unit to produce (a) a substantially solute free water stream and (b) a solute containing waste stream.

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109. (Original) The apparatus as set forth in claim 108 further including means for directing said solute containing waste stream to the inlet of said degasifier for further processing.

110. (Original) The apparatus as set forth in claim 105, further comprising de-oiling apparatus upstream of said multi-valent cation removal softener.

111. (Original) The apparatus as set forth in claim 103, further comprising filtration equipment downstream of said softener.

112. (Original) The apparatus as set forth in claim 103, further including means for directing backwash water to the inlet of said softener for further processing.

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REMARKS

The present Preliminary Amendment is being filed in order to provide an Abstract of the Disclosure as new page 44 of the specification, to make of record the claim to priority and to rewrite several claims to bring these claims into the proper format for U.S. prosecution.

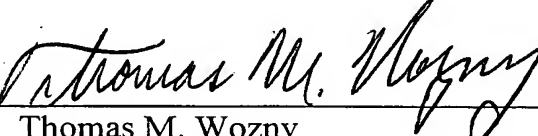
Applicant believes the application is in condition for examination and respectfully requests same.

Examination of this application is requested.

Respectfully submitted,

ANDRUS, SCEALES, STARKE & SAWALL, LLP

By



Thomas M. Wozny
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METHOD AND APPARATUS FOR HIGH EFFICIENCY EVAPORATION OPERATION

ABSTRACT

A process for treatment of an aqueous stream to produce a low solute containing distillate stream and a high solute/solids containing blowdown stream utilizing a method to increase the efficiency of an evaporator while providing an essentially scale free environment for the heat transfer surface. Multi-valent ions and non-hydroxide alkalinity are removed from aqueous feed streams to very low levels and then the pH is increased preferably to about 9 or higher to increase the ionization of low ionizable constituents in the aqueous solution. In this manner, species such as silica and boron become highly ionized, and their solubility in the concentrated solution that is present in the evaporation equipment is significantly increased. The result of this is high allowable concentration factors and a corresponding increase in the recovery of high quality reusable water with essentially no scaling.

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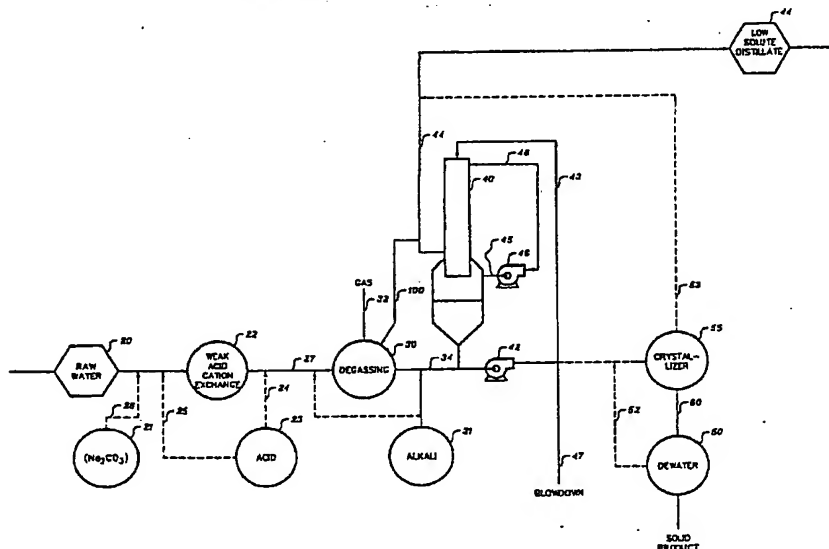
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METHOD AND APPARATUS FOR HIGH EFFICIENCY EVAPORATION OPERATION

CROSS-REFERENCE TO RELATED APPLICATIONS

- [00001] This application is based on and claims priority from provisional patent Application Number 60/419,552 filed on October 18, 2002 and provisional patent Application Number 60/419,549 filed on October 18, 2002.

TECHNICAL FIELD

- [00002] This process relates generally to a method and to a water evaporation system for the treatment of waters that contain dissolved organic materials and inorganic salts and in particular to a method that results in a non-scaling heat transfer surface. In various embodiments, this process relates to methods for feed water pretreatment that results in higher design concentration factors (higher recovery), an increase of the on-stream availability of the evaporation system, and an essentially scale free environment at the heat transfer surface.

BACKGROUND

- [00003] In today's world of increased awareness of the environment along with the high costs and regulations that prohibit and/or limit wastewater disposal to publicly owned treatment services or the environment, there is a demand for water treatment equipment that minimizes wastewater, promotes water reuse in the process, and lowers the quantity of fresh water that has to be imported from wells or public water supplies.

- [00004] The restraints put on many industries, such as steam-electric power plants, nuclear power plants, and oil production companies, have led to adoption of a Zero Liquid Discharge (ZLD) policy in many instances. A facility can achieve ZLD by collecting and recovering most or all of the water from the wastewater. The

resulting highly concentrated wastewater, or dry solids, are then held in ponds on site or the dry solids can be transported to a landfill.

[00005] A variety of technologies have been developed to recover water from wastewater or to reduce the volume of the wastewater. These technologies have limitations of complexity and susceptibility to interruption of service or failure of components due to corrosion, fouling, or scaling by the wastewater constituents, especially when feed waters vary from foreseen conditions.

[00006] A continuing demand exists for a simple and efficient process which can reliably provide water of a desired quality, in equipment that requires a minimum of maintenance. In particular, it would be desirable to improve efficiency of feed water usage, and lower both operating costs and capital costs for high quality water systems as is required for the various industries.

[00007] In most water treatment systems for the aforementioned industries, the plant design and operational parameters generally are tied to final concentrations (usually expressed as total dissolved solids, or "TDS"), which are tolerable in selected equipment with respect to the solubility limits of the sparingly soluble species present. In particular, silica, calcium sulfate, barium sulfate, calcium fluoride, and phosphate salts often limit final concentrations achievable or require operation of the system using the so-called seeded slurry design. To avoid scale formation and resulting decreases in heat throughput, the design and operation of an evaporation based water treatment plant must recognize the possibility of silica and other types of scale formation, and must limit water recovery rates and operational practices accordingly. In fact, typical evaporation plant experience has been that a reduction in distillate flow rates requires chemical cleaning of the evaporator at regular intervals. Such cleaning has been typically required because of scaling, particulate fouling, biofouling, or some combination thereof. Because of the cost, inconvenience, and production losses resulting from such cleaning cycles, it would be advantageous to lengthen the time between required chemical cleaning events as long as possible.

- [00008] It would be desirable to reduce the scaling, fouling, and corrosion tendencies of the feed water to the point where concentration factors could be increased in the design, and where flux rates could be increased, compared to limits of conventional scale control methods used in water evaporation systems. Raising the allowable concentration factors and flux rates, along with lowering the corrosion potential, is always important to the end user as these design points result in a lowering of capital costs.
- [00009] Present state of the art embodies several different strategies to alleviate the problems associated with scaling and fouling in higher concentration systems. These include the use of chelating agents, dispersants, solubility promoters, filters, silica precipitators, operating at low concentration factors, and the use of preferential deposition in a seeded slurry of calcium sulfate (CaSO_4) crystals. In the preferential deposition method, the low solubility precipitating crystals tend to deposit on the seeds that are suspended in the circulating solution rather than on the heat transfer surface.
- [00010] Membrane separation processes have also been used to obtain reusable water from wastewaters but they are typically limited to low recovery operations due to fouling/scaling limits, frequent cleanings, and replacement intervals of three years or less due, in part, to the frequent cleanings which can cause them to lose their rejection capability as well as productivity. A newly patented RO technology, HERO[™], utilizes softening and high pH operation to obtain recoveries up to 90 percent but has yet to show an extended membrane life comparable to the 20 years expected of an evaporator. This process is also limited in allowable concentration factor attainable due to osmotic pressure limitations, which currently is around about ten percent total dissolved solids.
- [00011] The prior art methods have the following shortcomings: (a) they rely on anti-scaling additives to prevent scale formation, or (b) they rely on seeding techniques for preferential deposition to minimize scaling of the heat transfer and other surfaces. Preferential deposition, while it works well in some applications, is not

the final answer as it cannot be expected to pick up every individual crystal that is precipitating and some invariably end up on the heat transfer surface, or sump walls, where they themselves then act as a seed site for scale buildup. In addition, certain feed waters do not have enough calcium sulfate (CaSO_4) in solution to serve as a self-renewing seed slurry. These feed waters then require the use of additional chemical treatment systems to supply the needed calcium (Ca) or sulfate (SO_4), or both, needed for this type of scale control method. Further complications inherent to the preferential deposition method are, (1) the need to carefully control the amount and size of seed that is circulating at any given time as too small a seed will cause fouling to occur in the laminar flow portion of the stream and too much seed will result in plugging of areas like water distribution trays, and (2) there is a limit to the concentration factor obtainable when the presence of double salts, such as glauberite ($\text{NaCa}(\text{SO}_4)_2$) will form scale as the concentration factor is increased.

[00012] Thus, for the most part, the prior art methods have one or more of the following shortcomings: (a) they rely on anti-scaling or dispersant additives to prevent scale formation, (b) are subject to scaling, fouling, and a short useful life, (c) they rely on seeding techniques to minimize scale deposition, or (d) are not able to concentrate beyond 7 or 8 percent TDS. Thus, the advantages of our treatment process, which exploits (a) multi-valent cation removal to non-precipitating residual levels, and (b) efficient dealkalization, to allow extended trouble free evaporator operation at high pH levels, are important and self-evident.

[00013] As water is becoming increasingly expensive, or in short supply, or both, it would be desirable to increase the ratio of treated product water to raw water feed in evaporator systems. Therefore, it can be appreciated that it would be desirable to achieve reduced costs of water treatment by enabling water treatment at higher overall concentration factors than is commonly achieved today. Finally, it would be clearly desirable to meet such increasingly difficult water treatment objectives with better system availability and longer run times than is commonly achieved today.

[00014] In so far as we are aware, no one heretofore has thought it feasible to operate an evaporator based water treatment system in a scale free environment and at an elevated pH, in continuous, sustainable, long-term operations to produce a high quality water product. The conventional engineering approach has been to design around or battle scale formation, by use of moderate pH, by limiting final concentration factors, by use of chemical additives, or by use of preferential deposition.

[00015] In contrast to prior art methods for water treatment, the method described herein uses the essential design philosophy of virtually eliminating any possible occurrence of scaling phenomenon during evaporator operation at the maximum feasible pH, while maintaining the desired concentration factor, and taking the benefit of water recovery that results.

SUMMARY

[00016] We have now invented a novel water treatment method that emphasizes feeding an evaporator with an essentially multi-valent cation free water that allows high pH evaporation in a scale free environment, to produce a high quality distillate at greater cycles of concentration.

[00017] In a unique feed water treatment process, raw feed waters of suitable chemical composition are preferentially treated with a weak acid cation ion exchange resin, operated in the hydrogen form, to simultaneously remove multi-valent cations and alkalinity. The weak acid cation ion exchange resins can be operated at incoming raw feed water hardness and alkalinity levels well above those that would cause conventional ion exchange systems to fail due to hardness breakthrough.

[00018] The preferred treatment train design used in our wastewater treatment plant overcomes a number of important and serious problems. First, the low levels of multi-valent cations, combined with virtual elimination of non-hydroxide alkalinity, substantially eliminates the precipitation of scale forming compounds associated

with sulfate, carbonate, or silicate anions. Thus, cleaning requirements are minimized. This is important commercially because it enables a water treatment plant to avoid lost water production, which would otherwise undesirably require increased treatment plant size to accommodate for the lost production during cleaning cycles. Second, the preferred high pH operational conditions enable a high degree of ionization to be achieved in various species which are sparingly ionized at neutral or near neutral pH in aqueous solution, to enable such species to be concentrated to higher levels before precipitation. Third, the method does not have the osmotic pressure limitation of membrane based systems and allows operation and much higher TDS concentrations with resultant higher recovery or water and reduction in final waste quantity. Finally, operation at high pH and heat provides protection against biological contamination, thus preventing undesirable contamination of the distillate stream. At the preferred high operational pH, bacteria and endotoxins are effectively destroyed. In essence, water treatment systems operated according to the process herein normally operate at conditions, which might ordinarily be considered cleaning conditions for conventional evaporation systems.

[00019] We have now developed a novel process design for use in the treatment of water. In one embodiment, the process involves treatment of a feed water stream, which is characterized by the presence of (i) multi-valent cations, (ii) alkalinity, and (iii) molecular species which are sparingly ionized when in neutral or near neutral pH aqueous solutions, to produce a low solute containing distillate stream and a high solids containing blowdown stream. The process involves effectively eliminating the tendency of the raw feed water to form scale when the raw feed water is concentrated to the desired concentration factor at a selected pH by removing multi-valent cations from the raw feed water stream and by effecting, in any order, one or more of the following: , (i) removing alkalinity from the raw feed water stream, (ii) removing dissolved gases whether initially present or created during the multi-valent cation or alkalinity removal steps, or (iii) raising the pH.

The pH of the feed water is raised to a selected pH in a range between 9 and 10, or otherwise in excess of 10, and more preferably to about 11 to 12 or somewhat more, until the benefits gained by high ionization of silica and other species is outweighed by the additional cost. The pH increase is accomplished by adding a selected base to the softened and degassed feed stream, preferably by direct injection into the feed stream or alternately into the sump of the evaporator. The pH increase urges the molecular species, which are sparingly ionized when in neutral or near neutral pH toward increased ionization. The pH adjusted feed water is then sent through heat transfer evaporation equipment to produce a concentrated blowdown stream and a low solute containing distillate stream. The evaporation equipment is typically of the falling film type wherein the heat transfer surface is comprised of a number of tubes with evaporation on either the interior or exterior surface, a plurality of plate style with evaporation on the outer surface, or a forced circulation process. It is important that in our process, the evaporation equipment operates in an essentially scale free environment to produce a distillate stream, which is substantially free of the normally undesirable species while operating at an increased efficiency due to increased solubility limits of sparingly soluble salts at an elevated pH.

OBJECTS, ADVANTAGES, AND FEATURES

[00020] From the foregoing, it will be apparent that one important and primary object of the present invention resides in the provision of a novel method for treatment of water to reliably and continuously produce, over long operational cycles, a water distillate stream of high quality, suitable for reuse, at a reduced capital and operating cost.

[00021] More specifically, an important object of our invention is to provide an evaporation based water treatment method which is capable of avoiding common scaling and fouling problems, so as to reliably provide a method of high quality

water generation when operating at increased efficiency on a variety of wastewaters.

- [00022] Other important but more specific objects of the invention reside in the provision of a method for water treatment as described in the preceding paragraphs which:
- [00023] allows for the removal of multi-valent cations and alkalinity from a selected feed water to be done in a simple, direct manner;
- [00024] has high efficiency rates, that is, provide high product water outputs relative to the quantity of feed water input to the water treatment plant;
- [00025] allows operation at pH above 9, which reduces the concentration of hydrogen ion present in the aqueous solution;
- [00026] allows operation at higher specific heat transfer rates, which reduces the amount of heat transfer surface required;
- [00027] allows removal of dissolved oxygen from the aqueous solution;
- [00028] in conjunction with the preceding objects, the reduction of hydrogen ion and oxygen concentration reduces the corrosiveness of the aqueous solution allowing the use of lower cost materials for most feed waters;
- [00029] provide lower unit costs to the water treatment plant operator and thus to the water user, than is presently the case;
- [00030] in conjunction with the just mentioned object, results in less chemical usage than in most water treatment facilities, by virtually eliminating use of some types of heretofore commonly used chemical additives, particularly scale inhibitors or chemicals needed to maintain a seeded slurry, and eliminates expensive physical/chemical scale removal techniques and downtime;
- [00031] in conjunction with the scale free environment object previously mentioned, results in a lower corrosion potential and allows for lower grade materials of construction in lieu of high alloy materials.

[00032] A feature of one embodiment of the present invention is the use of a unique combination of weak acid cation ion exchange with substantially complete hardness and alkalinity removal, and subsequent high pH evaporation operation, thereby enabling the water treatment plant to minimize the percentage of blowdown water. This results in high overall cycle efficiencies.

[00033] Another feature of the present invention is the use of a high pH operation to highly ionize weakly ionizable species such as silica or boron, thus enabling operation with silica or boron concentration limits considerably exceeding the limits of conventional evaporation treatment systems when treating feed waters of comparable chemistry.

[00034] Another feature of the present invention is the capability to remove ammonia from the feed stream as a part of the process instead of another separate process. The ammonium ion (NH_4) is very soluble in water with a dissociation constant (pK_a) value of 9.24. At a pH of 11.2 in the feed stream, a typical process operating point of the present invention, it is over ninety nine percent (99%) dissociated into the ammonia (NH_3) ion and can be removed in the degassifier.

[00035] Yet, another feature of the present invention is the capability to retrofit existing evaporation plants to operate according to the present process design, to increase capacity without increasing the installed heat transfer surface.

[00036] Other important objects, features, and additional advantages of the invention will become apparent to those skilled in the art from the foregoing, and from the detailed description which follows, and from the appended claims, in conjunction with the accompanying drawings.

BRIEF DESCRIPTION OF THE DRAWINGS

[00037] All the exemplary embodiments shown herein incorporate the Zero Liquid Discharge (ZLD) concept option as a part of the illustration. Those skilled in the art will recognize that merely minimizing the blowdown stream without the use of a dewatering device may, on certain occasions, also qualify the system as ZLD.

The high efficiency evaporation method is highly site specific wherein individual process steps are customized to fit the specific feed water, and needs of the customer, at the specific site. For that reason, all possible embodiments of this novel method of water treatment are not illustrated and, as those skilled in the art can appreciate, other illustrative embodiments would merely reflect variations and arrangement of some components without affecting the spirit or concept of this invention.

- [00038] The same identifier will reference identical features depicted in each of the drawings.
- [00039] Figure 1 is a graph illustrating the ionization of silica as a function of pH;
- [00040] Figure 2 is a flow diagram illustrating one embodiment of the novel water treatment method disclosed herein to obtain high efficiency evaporation utilizing a weak acid cation exchange system to remove divalent cations and alkalinity associated with hardness in one step;
- [00041] Figure 3 is a flow diagram illustrating another embodiment of the novel water treatment method disclosed herein to obtain high efficiency evaporation and high purity distillate simultaneously;
- [00042] Figure 4 is a flow diagram illustrating another embodiment of the novel water treatment method disclosed herein to obtain high efficiency evaporation, minimized blowdown, and low solute containing distillate for use as cooling tower or scrubber makeup;
- [00043] Figure 5 is a flow diagram illustrating another embodiment of the novel water treatment method disclosed herein showing the arrangement of equipment wherein sodium zeolite softening is sufficient for high efficiency evaporation. The use of an optional lime or lime/soda softener for hardness removal is also depicted; and
- [00044] Figure 6 is a flow diagram illustrating another embodiment of the novel water treatment method disclosed herein wherein acid addition adequately removes

any alkalinity present in the feed stream and where hardness removal, if present, can be optionally accomplished with lime or lime/soda softening.

DETAILED DESCRIPTION

[00045] Since many industrial applications of various types generate large quantities of wastewater that is becoming increasingly expensive and regulated, it has become desirable to process it for internal reuse and limit or eliminate discharge into public utilities. Present day state of the art is limited on how much water can be recovered by scale causing ions, such as hardness and silica, which are inherent in these waste streams. The addition of expensive scale inhibiting agents or scale control methods are beneficial but still have their limits of usefulness. We have designed a novel process to overcome these limitations and recover more of the wastewater for reuse than was previously possible by providing a scale free environment in the evaporator. As used herein the term "scale" is intended to encompass not only a thin coating, layer or incrustation (usually rich in sulfate or calcium) that is deposited on a surface, but also particulate fouling, biological fouling, or some combination thereof.

[00046] Attributes that characterize the high efficiency evaporator (HEVAP) process design and operation are:

- [00047] (1) Very high solubility of weak acid anions such as silica.
- [00048] (2) Very high achievable concentration factors (recovery--ninety percent (99%) or higher recovery can be achieved).
- [00049] (3) Biological fouling is essentially eliminated.
- [00050] (4) Particulate fouling is substantially reduced.
- [00051] (5) Cleaning frequency is substantially reduced.
- [00052] (6) Addition of scale inhibitors is virtually eliminated.
- [00053] (7) Corrosion potential is reduced.
- [00054] (8) Higher heat flux is achievable.

- [00055] (9) Reduced overall capital cost, compared to conventional evaporation systems.
- [00056] (10) Reduced overall operating cost, compared to conventional evaporation systems.
- [00057] The HEVAP evaporation system is highly site-specific. Individual process steps are customized to fit the specific feed water at a specific site. Regardless of the difference in the pretreatment process for different sites, one process parameter is common for all applications, namely that the evaporator system is operated at the highest feasible blowdown pH and that the circulating solution provides a scale free environment at the heat transfer surface.
- [00058] In order to operate an evaporative system with a pH of at least about 9.0, preferably at least about 10.5, and most preferably between 11 and 12, or above, several process conditions must be met in order to effectively eliminate the potential for scale formation on the heat transfer surface. Some of those process conditions are also necessary for operating an evaporative system at very high concentration factors. Such process conditions are as follows:
- [00059] (1) The calcium, magnesium, strontium, and barium concentration in the evaporator feed must be substantially absent, preferably to near zero, and most preferably, to essentially zero.
- [00060] (2) Aluminum, iron, manganese, and other multi-valent cation content including organically bound species, as well as the presence of colloidal particles containing such materials, should be substantially absent, and preferably to near zero.
- [00061] (3) Buffering anions (specifically bicarbonate, or carbonate, and/or phosphate species) should be reduced to as low of a level as can be practically achieved.
- [00062] (4) Dissolved and suspended gasses such as oxygen, ammonia, and others should be minimized.

[00063] The selection of specific operations and control points, to fulfill the above process condition requirements, is influenced by the characteristics of each specific feed water. The concentration factor needed (or desired for a specific application) also affects the operations and control point criteria as well. FIG. 2 represents a highly effective evaporator unit process sequence.

[00064] The first step is to adjust the hardness-to-alkalinity ratio of the feed water, if needed. Optimizing this ratio, which is typically done by alkali addition, makes complete hardness removal feasible in a weak acid cation ion exchange process operated in the hydrogen form as described in the next process step.

[00065] The second step in the evaporator process train involves the utilization of a weak acid cation (WAC) resin (e.g. DOWEX.RTM. MAC-3, or Lewatit CNP-80, Amberlite.RTM. IRC-86). Operated in either the hydrogen or sodium form, the WAC resins remove multi-valent cations and, in the hydrogen form removes any alkalinity associated with hardness.

[00066] The third step involves pH adjustment by adding acid to the WAC effluent. Acid is added to destroy any alkalinity remaining, after multi-valent cation removal, if any such alkalinity is present.

[00067] In a fourth step, the acidified effluent, containing virtually zero alkalinity, is then treated for carbon dioxide removal. This removal can be accomplished in any of various type degasifiers. The degasified feed water stream with multi-valent cation levels below the limits required for scale free operation and essentially zero alkalinity, is then injected with a soluble alkali, preferably for adjusting pH to 9.0 or higher, more preferably 10.5 or higher, and most preferably 11.0 or above.

[00068] In other embodiments where ammonia is a concern, a variation of the steps is required. Alkali is added to the feed stream to decrease ammonia solubility at elevated pH prior to removing it as a gas in a degassifier. In applications where both alkalinity and ammonia are present, two degassifiers are required, one for removing any gases such as carbon dioxide at low pH and one for ammonia removal at high pH. In cases where there is no alkalinity due to a very low pH in

the feed stream, the ammonia can be removed by injecting alkali prior to a single degassifier after reducing the multi-valent cations to a non-scaling level in the feed stream.

[00069] Feed waters utilized for production of reusable water, especially those encountered in wastewater treatment, include the presence of silicon dioxide (also known as silica or SiO_2) in one form or another, depending upon pH and the other species present in the water. For evaporator systems, scaling of the heat transfer surface with silica is to be religiously avoided. This is because (a) silica forms a relatively hard scale that reduces productivity of the evaporator, (b) is usually rather difficult to remove, (c) the scale removal process produces undesirable quantities of spent cleaning chemicals, and (d) cleaning cycles result in undesirable and unproductive off-line periods for the equipment. Therefore, regardless of the level of silica in the incoming raw feed water, operation of conventional evaporation processes, without a preferential deposition seeded slurry process, generally involves concentration of SiO_2 in the high solids stream to a level not appreciably in excess of 150 ppm of SiO_2 (as SiO_2). This requires that evaporator systems be operated at lowered concentration factors (recovery rates) to prevent silica concentration in the blowdown stream from exceeding solubility limits. Seeded slurry systems can be taken to concentration factors that surpass the solubility of silica but rely on seed management procedures and are still prone to scaling of the evaporator.

[00070] Scaling due to various scale forming compounds, such as calcium sulfate, calcium carbonate, and the like, can be predicted by those of ordinary skill in the art and to whom this specification is directed, by use of the Langelier Saturation Index (LSI) or the Stiff-Davis Index (S&DI), or other available solubility data. Operating parameters, including temperature, pH, distillate and blowdown flow rates, must be properly accounted for, as well as the various species of ions in the raw feed water, and those species added during pretreatment. The Nalco Water

Handbook, copyright 1979, by McGraw-Hill details the procedure for use of the indexes.

[00071] With reference to Figure 2, wastewater stream 20 of this invention will typically contain, soluble and insoluble, organic and inorganic components. The inorganic components can be salts such as sodium chloride, sodium sulfate, calcium chloride, calcium carbonate, calcium phosphate, barium chloride, barium sulfate, and other like compounds. Metals such as copper, nickel, lead, zinc, arsenic, iron, cobalt, cadmium, strontium, magnesium, boron, chromium, and the like may also be included. When treating a wastewater stream from an oil refinery, organic components will be present and are typically dissolved and emulsified hydrocarbons such as benzene, toluene, phenol, and the like.

[00072] It is commonly understood that the solubility of silica increases with increasing pH, and that silica is quite soluble in high pH aqueous solution. Along with solubility, the degree of ionization of silica also increases with increasing pH. While the increase in silica solubility is not directly proportional to the degree of ionization, the rate of increase in silica solubility is basically proportional to the rate of change in ionization as increased ionization results in the soluble silicate ion being the dominant species. This discrepancy between solubility and ionization is explained by the fact that even undissociated silica exhibits some solubility in aqueous solutions, typically up to about one hundred twenty (120) ppm to one hundred sixty (160) ppm, depending upon temperature and other factors. In comparison, it has been demonstrated that silica solubility at pH 10.5 is in excess of one thousand five hundred (1,500) ppm at ambient temperature; silica is increasingly soluble as temperature and/or pH increases.

[00073] Silica is very weakly ionized when in neutral or near neutral aqueous solutions and is generally considered to exist as undissociated (meta/ortho-) silicic acid ($H_4 SiO_4$) in most naturally occurring waters with a pH of up to about 8. The dissociation constant (pKa) value for the first stage of dissociation of silica has been reported at approximately 9.7, which indicates that silica is approximately

fifty percent (50%) ionized at a pH of 9.7; the other fifty percent (50%) remains as undissociated (ortho) silicic acid at that pH. A graphical representation of the relationship between pH and the percent silica ionization is shown in FIG. 1.

Clearly, it would be advantageous, where silica ionization is desired, to operate at a pH in excess of 10, and more preferably, in excess of 11, and yet more preferably, in excess of 12 where all of the silica molecule is present as a soluble silicate ion.

[00074] Therefore, increasing the pH of the evaporator operation thus provides the major benefit of increased silica solubility. To gain maximum benefit from silica ionization at high pH, the evaporator system should be operated at a pH as high as possible. Preferably, the evaporator system is operated at a pH of about 10.5 or above, and more preferably, at a pH of 11 or higher. This contrasts with typical evaporator operation practice, where operating pH has been maintained at less than 9 in order to avoid scale formation, particularly silica and carbonate scales.

[00075] Referring to FIG. 2, one embodiment of this process for evaporation equipment operation is shown. In this method, raw water 20 is first treated in a weak acid cation (WAC) ion exchange unit 22, where hardness and bicarbonate alkalinity are simultaneously removed. For those cases where raw water 20 hardness is greater than alkalinity, operation of the weak acid cation ion exchange unit 22 must be facilitated by addition of a source of alkalinity 21, such as by addition of an aqueous solution of sodium carbonate (Na_2CO_3). Preferably, the WAC unit 22 is operated in the hydrogen form for ease of operation and regeneration. However, it would also work in the sodium form, followed by acid addition. In any case, in the just mentioned case and otherwise optionally where appropriate, acid 23 is added to the effluent 27 from the WAC unit(s) 22 to enhance bicarbonate destruction. Sufficient acid is added to lower the pH where bound carbonates are converted to a free gas carbon dioxide. Then, the carbon dioxide 32 that has been created in the WAC (and/or by acid addition) is removed, along with other non-condensable gasses such as oxygen and nitrogen, preferably in an atmospheric pressure or vacuum/flash degasifier 30. Finally, an alkali 31 (base) is

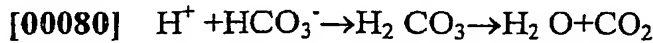
added, preferably by pumped injection of liquid solution, to increase the pH of the feed water 34 to a selected level. Any of a variety of conveniently available and cost effective base products may be used, provided that no appreciable scaling tendency is introduced. Besides use of common sodium hydroxide, other chemicals such as sodium carbonate, potassium hydroxide, or potassium carbonate might be selected. In fact, in certain cases, an organic base, such as a pyridine type compound, may be used effectively to carry out this process.

[00076] The pH of the feed water is raised to a selected pH of at least about 9.0, or up to about 10, or preferably to a range between 10 and 11, or otherwise in excess of 11, and more preferably to 12 or more, and most preferably, to 13 or more.

[00077] The weak acid cation ("WAC") ion-exchange resins used in the first step of the preferred embodiment of the method defined herein, as illustrated in FIG. 2, are quite efficient in the removal of hardness associated with alkalinity. Such a reaction proceeds as follows:



[00079] Then, the hydrogen combines with the bicarbonate to form carbonic acid, which when depressurized, forms water and carbon dioxide, as follows:



[00081] Regeneration of the resin is accomplished by use of conveniently available and cost effective acid. It is well known by those in the art that regeneration of WAC ion-exchange resins may proceed quite efficiently, at near stoichiometric levels (generally, not more than about one hundred and twenty percent (120%) of ideal levels). Preferably, hydrochloric acid may be used, since in such cases highly soluble calcium chloride would be produced, and the regeneration process would not pose the potential danger of formation of insoluble sulfate precipitates, such as

calcium sulfate, even with high strength acids. However, by use of a staged regeneration procedure, i.e., by using a low concentration acid followed by a higher concentration acid, it is possible to reliably utilize other acids, including sulfuric acid ($H_2 SO_4$), while still avoiding undesirable precipitates on the resin. In this manner, hardness ions are solubilized to form soluble salts, which are eluted from the resin bed and are typically sewered.

[00082] Other polyvalent cations, most commonly iron (Fe^{++}/Fe^{+++}), magnesium (Mg^{++}), barium (Ba^{++}), strontium (Sr^{++}), aluminum (Al^{+++}), and manganese (Mn^{++}/Mn^{++++}), are also removed by the WAC resin. Since the presence of even very small quantities of hardness or other species of decreasing solubility at increasing pH will result in precipitation of sparingly soluble salts under the process conditions present in our process, care must be taken to prevent precipitation on the heat transfer surface of the substances such as calcium carbonate, calcium hydroxide, magnesium hydroxide, and magnesium silicate. One precaution that should be observed is that both hardness and non-hydroxide forms of alkalinity should be aggressively reduced in the feed water, prior to upward pH adjustment to selected evaporator operating conditions. Once the multi-valent cations and non-hydroxide forms of alkalinity have been removed, then the desired pH increase may be accomplished with any convenient alkali source, such as sodium or potassium alkali. Once this pretreatment has been thoroughly accomplished, then an evaporator system can be safely operated at very high pH levels, in order to take advantage of the aforementioned silica solubility.

[00083] The treated and conditioned feed water 34 is directed into the evaporator 40 where it mixes with and dilutes the concentrated high solids stream 43. This stream is recirculated with pump 42 and a small portion is removed as evaporator blowdown 47 on each pass through the evaporator 40. In the evaporator 40 the solutes in the feed water 34 are concentrated by removing water from the diluted recirculating solution 43 as it passes over the heat transfer surface. As depicted in Figure 2, the evaporator utilizes falling thin film evaporation wherein the

recirculated stream 43 is thinly spread across the inner surface of a plurality of heat transfer tubes. A small portion of water is removed from the thin recirculating stream in the form of steam 45 driven by heated, compressed steam 48 which is condensing on the outside of the heat transfer tubes. The water that has been removed, in the form of steam 45, is compressed through the compressor 46, and the compressed steam 48 is condensed on the outer surface of the heat transfer tubes to generate more steam 45, and keep the evaporation process going. The condensing steam 48 is known as distillate or condensate, as is known to those skilled in the art of evaporation, and contains a low level of non-volatile solutes, typically, in some embodiments, less than 10 parts per million (ppm). It should be noted that the use of a tubular falling film evaporator 40 design is provided only for purposes of enabling the reader to understand the evaporation process and is not intended to limit the process to the use of the same. Those familiar with the art will recognize that other designs, such as, for example, a rising film evaporator, or a forced circulation evaporator, or a plate style evaporator may be alternately utilized with the accompanying benefits and/or drawbacks that may be inherent in the alternative designs.

- [00084] The condensing steam 48 descends by gravity to the bottom of the tubular heat transfer surface and is collected as distillate stream 44. A small portion of the distillate 44 may be sent to the earlier discussed degasser 30 via line 100 for use in mass transfer, i.e., adding heat to the feed water stream 27 to remove non-condensable gasses such as carbon dioxide 32. However, the bulk of the distillate 44 is directed to the terminal point of the evaporator where it is available for use in any process that requires high quality water as a makeup stream. Typical, but not limiting, uses include those shown in the Figure 4 embodiment of the high efficiency evaporation process in which the distillate can be used as makeup to a cooling tower or scrubber. Other uses would include low-pressure boilers and, in the hydrocarbon recovery field (produced water), as feed water to a once through

steam generator (OTSG) that generates steam for injection into oil-bearing formations.

[00085] Although the low solute containing distillate 44 produced by the evaporator is relatively pure water, there are instances where a higher purity is required. Figure 3 depicts an embodiment wherein several different options are shown for obtaining different levels of high purity. In most cases the residual solutes in the distillate stream 44 involve salts other than hardness. In one embodiment, the distillate 44 is passed through a cation ion exchange system 70, followed by an anion ion exchange system 72, and then polished in a mixed bed ion exchange system 76 to produce a very high purity water. The inclusion of all three ion exchange systems is for illustration only and those of ordinary skill in the ion exchange arts and to which this disclosure is directed will recognize that only those ion exchange systems required to meet the requirements for purity will be used. In any event, the ion exchange systems will require regenerant chemicals and that will result in a regeneration waste stream 73 which can be directed to the inlet of the degasifier 30 for further treatment in the evaporator 40. In an alternate embodiment, the removal of residual solutes in the distillate stream 44 can be accomplished by passing the stream through an electrodeionization (EDI) unit 80. The EDI reject is also capable of being recycled to the evaporator by directing it to the inlet of the degasifier 30.

[00086] The impact of very low levels of silica, etc., in the relatively pure distillate 44 obtainable by evaporation on the behavior/operation of a post-evaporator ion exchange system is extremely significant. Since the vast majority of post-evaporator ion exchange is regenerated on the basis of either silica or boron breakthrough, a factor of ten reduction in the influent silica/boron content will provide much longer run times between regenerations. Absence of carbon dioxide, as well as bicarbonate in the distillate 44 (due to a high pH, typically at least 10), will also increase on-line time before silica/boron leakage exceeds normal threshold values. Reduction of strongly ionized species concentration in the distillate 44 is of

relatively less significance, since most post-evaporator ion exchange is ultimately silica or boron limited.

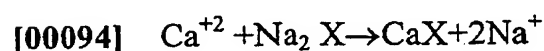
- [00087] The just described combination of treatment steps when combined with the novel process described hereinabove produces a water of sufficient quality, and economic quantity, to be used in high pressure and packaged boilers. Typical users would be the power generation industry and hydrocarbon recovery operations wherein 100% quality steam is utilized for steam flooding applications.
- [00088] The evaporator blowdown 47 containing the concentrated solutes originally present in the feed water 34 along with any chemicals used to raise the pH and/or regenerate post ion exchange systems can be disposed of by the standard approach used at individual sites. This includes holding on-site in waste evaporation ponds, trucking to a waste site, or injection into deep wells.
- [00089] Alternatively, the blowdown stream 47 can be directed to a crystallizer 55 that further processes the concentrated stream to recover low solute distillate 53 and a high-suspended solids stream 60. The distillate stream 53 is then combined with the falling film evaporator distillate stream 44 to effect increased recovery of the evaporator feed stream 34. The high-suspended solids containing stream 60 can then be directed to a dewatering device 50, typically a belt filter but alternatively a filter press or even a spray drier. The final product is a dried solid that is suitable for landfill or possibly even reused within the originating process. The two different filter methods generate a high solute/low suspended solids stream 52 that is directed back to the crystallizer 55 for further processing.
- [00090] In other embodiments, and as suited to meet the particularized needs of a selected raw feed water chemistry, various forms of hardness removal may be utilized, including sodium form strong acid cation exchange 65, followed by acidification (see FIG. 5) or even the use of a lime 82 (or similar lime/soda) softener as an additional pretreatment step to either sodium form strong acid cation exchange 65 or weak acid cation exchange 22 (see FIGS. 2 and 5). The direct injection of lime and sodium carbonate into the feed stream can also be utilized and

the resulting precipitate filtered out in a membrane separation process, such as ultrafiltration, as a substitute for the lime/lime soda 82 softener.

[00091] For particularly soft waters, the lime or lime/soda softener 82 may be totally inappropriate, and this method may proceed with no softening of the raw water, and only a simple acid 24 feed before degasifying, as can be seen in FIG. 6. On the other hand, where softening is appropriate, some raw feed waters can be appropriately treated for reductions in hardness and alkalinity to a desired extent by softener 82.

[00092] In still other embodiments and for a selected feed water, the use of softening membranes for partial removal of hardness may be incorporated into the process as a replacement for sodium zeolite 65 or weak acid cation 22 softening.

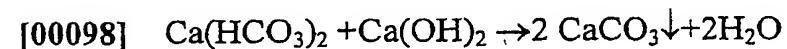
[00093] In cases where raw water composition is such that sodium zeolite softening is advantageous, as is depicted in FIG. 5, elimination of calcium hardness proceeds as follows:

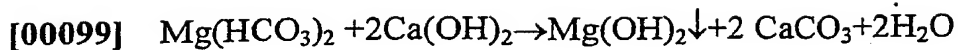


[00095] Then, bicarbonate alkalinity is converted to carbon dioxide, with a selected acid, in a manner similar to the following:



[00097] For those waters where lime softening may be an acceptable or preferred method for initial hardness and alkalinity reduction, the addition of lime to the water reduces calcium and magnesium hardness, and associated bicarbonate alkalinity, as follows:





[000100] Regardless of the equipment configuration selected for treatment of a particular raw water chemistry, the key process parameters are (a) to remove those cations which, in combination with other species present at high pH, would tend to precipitate sparingly soluble salts on the heat transfer surfaces, and (b) eliminate non-hydroxide alkalinity to the maximum extent feasible, to further protect against precipitation of scales on the heat transfer surfaces when operating at an elevated pH.

[000101] FIG. 10 illustrates the use of our novel method of evaporator system operation for cooling tower makeup water or for scrubber makeup water. The evaporator unit 40 and various pretreatment equipment are operated according to the methods set forth hereinabove, to produce a high quality distillate 44. Although the cooling tower 95 and scrubber 90 could be fed with distillate 44, more typically, the cooling tower 95 and scrubber 90, for example in a steam-electric power plant, would be supplied by usual raw water 20 supplies, such as municipal or well water. Therefore, cooling tower blowdown 96 and scrubber blowdown 91 are typically high in both hardness and alkalinity. Likewise, this system may be used to treat water having intimate contact with ash, such as ash pond water or ash-sluicing water from coal fired steam-electric power plants. In our evaporation process, a significant amount of reusable water can usually be obtained by our method of evaporation pretreatment and operation, unlike the case with conventional evaporative systems.

[000102] Another advantage, since an evaporator system when operated as described herein will not be subject to scaling or fouling conditions, wastewaters from refineries, hydrocarbon recovery operations, pulping and papermaking operations, membrane concentration systems, and municipal sewage treatment plants, are candidates as suppliers of raw water 20. Typical industrial uses where water of

sufficient quality may be attained when treating wastewaters include cooling towers, boiler makeup, scrubber makeup, and the like.

[000103] Benefits of HEVAP Evaporation Process Design and Operation

[000104] Many exemplary and desirable process benefits provided by the HEVAP evaporation system process design and operation were listed above. Detailed explanation of such benefits include:

[000105] (A) High solubility of silica

[000106] It has been documented by others that silica solubility in water at 25°C. approaches 6000 milligram per liter (mg/l) at a pH of 11 and at a pH of 12, the solubility approaches 60,000 mg/l when in equilibrium with amorphous silica. It has also been documented that the solubility of silica in water goes up with an increase in temperature leading to the conclusion that evaporator operation at temperatures in excess of 100°C. at an elevated pH and silica levels up to 6000 mg/l is feasible. However, this is only possible if precipitating species such as calcium and magnesium and the like have been removed from the feed stream so that they cannot encourage the polymerization of silica and subsequent scaling on the heat transfer surface. The novel process disclosed herein wherein an aggressive approach to multi-valent cation and alkalinity removal is practiced, allows operation at much higher levels of silica concentration than were previously possible in normal evaporators. Since the high pH utilized by this novel process assures increased silica solubility, a concentration factor (i.e., ratio of feed rate 34 to blowdown rate 47) for the evaporator 40 can be selected so that silica solubility is not exceeded.

[000107] (B) High Recovery Rates

[000108] Since multi-valent ions such as calcium, magnesium, barium, strontium, aluminum, iron, manganese, etc., have been removed prior to concentration in the evaporator, undesirable precipitation of species such as calcium carbonate, calcium fluoride, calcium sulfate, barium sulfate, magnesium hydroxide, aluminum/magnesium silicate, etc., does not occur in the high efficiency evaporator process, and thus that type of precipitation no longer limits the recovery achievable by an evaporator system. Importantly, silica solubility is increased dramatically at the normal high efficiency brand evaporator operating pH (preferably at approximately 11 or above). Since silica usually represents the ultimate limiting criterion, in terms of maximum allowable concentration in an evaporator system, increased silica solubility along with essentially total absence of species such as calcium, barium, etc., in the evaporator feed, will allow evaporator operation at very high recovery rates (98 to greater than 99 percent) with the vast majority of feed waters.

[000109] (C) Biological Fouling Eliminated

[000110] Most commonly occurring microbial species are completely lysed (physically destroyed by wall rupture) at the high operating pH. In fact, even virus, spores, and endotoxins are either destroyed or rendered incapable of reproduction/proliferation at very high pH levels. Saponification of lipids (fat) is expected to play a role in the process as well since fatty acids and their corresponding glycerides will form soluble 'soaps' at the high operating pH. This characteristic of the new process can be of significant benefit for sites with known biofouling problems or for the treatment of bio-contaminated/bio-active wastewater.

- [000111] (D) Cleaning Frequency Reduced
- [000112] The HEVAP process, which utilizes aggressive removal of multi-valent cations and alkalinity along with a high pH in the evaporator, lengthens the time between shutdowns to clean the equipment. Typically, two weeks per year are used to clean heat transfer surfaces and sumps by opening them up to allow access for expensive high pressure hydro-blasting procedures to remove the bulk scaling material. This is then followed by time-consuming washes with costly proprietary chemicals to remove any scale not removed by hydro blasting.
- [000113] In contrast, the HEVAP process, by removing essentially all minimal solubility ions and alkalinity, incurs only minimal scaling due to small leakage from the softeners used. The result of this is that cleaning intervals can be extended and that they can be simply and effectively accomplished by commodity cleaning chemicals, such as hydrochloric acid solutions, tetra-sodium EDTA, and sodium hydroxide. Expensive proprietary chemical cleaning agents are not required. The scales that could occur would be predominantly calcium carbonate, magnesium hydroxide, magnesium silicate, and the like, all of which can be removed with a simple acid wash.
- [000114] The increased system availability, with minimal scaling and virtually non-existent bio-fouling, is clearly another important benefit of this novel operational method.
- [000115] (E) Scale Inhibitors
- [000116] The use of antiscalants, scale dispersants, scale inhibitors, or scale control methods, while not harmful or incompatible with the new process, can be minimized, if not completely eliminated, due to the aggressive removal of multi-valent cations along with virtually all non-hydroxide alkalinity as practiced by the HEVAP pretreatment process.

[000117] (F) Higher Flux

[000118] Present day state of the art evaporators are heat flux (flow) limited due to the presence of low solubility scale causing ions such as calcium carbonate, calcium sulfate, silica and the like. A higher flux can be incorporated into the evaporator design when these ions are absent in the feed stream and that is what is accomplished with the process described herein.

[000119] (G) Reduced Capital Cost

[000120] The lowered corrosion potential that results from operating the evaporator with a high pH in the concentrated circulating solution allows the use of lower cost materials for heat transfer tubes or plates and other wetted surfaces that are contacted by the concentrated solution, such as sump walls. This is an important advantage since the costs of these materials have a substantial impact on the capital cost of an evaporator. In most cases the use of high cost duplex and AL6XN (6 percent minimum molybdenum) type stainless steels, which are normally used in high chloride salt solutions, can be eliminated in favor of a lower grade stainless such as 316.

[000121] (H) Reduced Operating Cost

[000122] Water plant operating costs can be reduced due to minimizing, or eliminating, costly proprietary antiscalants and/or dispersants. Additional savings can be found by eliminating the need for seeded slurry operation at installations where the multi-valent ions are at a low level in the feed stream but are accompanied by high silica levels. Along with the cost of seeding the evaporator with calcium sulfate crystals, there is also incurred costs associated with calcium chloride and/or sodium sulfate injected chemicals to provide enough precipitating ions to maintain the seed bed at many installations. Further savings can be realized by the reduction in frequency of cleaning operations, less expensive cleaning chemicals, less downtime for cleaning, and no requirement for costly physical

cleaning operations. Still further, if the ZLD option is incorporated, the cost of sending the blowdown to a public utility company is eliminated.

[000123] It will thus be seen that the objects set forth above, including those made apparent from the preceding description, are efficiently attained, and, since certain changes may be made in carrying out the above method and in construction of a suitable apparatus in which to practice the method and in which to produce the desired product as set forth herein, it is to be understood that the invention may be embodied in other specific forms without departing from the spirit or essential characteristics thereof. For example, while we have set forth an exemplary design for simultaneous hardness and alkalinity removal, other embodiments are also feasible to attain the result of the principles of the method disclosed herein. Therefore, it will be understood that the foregoing description of representative embodiments of the invention have been presented only for purposes of illustration and for providing an understanding of the invention, and it is not intended to be exhaustive or restrictive, or to limit the invention to the precise forms disclosed. On the contrary, the intention is to cover all modifications, equivalents, and alternatives falling within the spirit and scope of the invention as expressed in the appended claims. As such, the claims are intended to cover the methods and structures described therein, and not only the equivalents or structural equivalents thereof, but also equivalent structures or methods. Thus, the scope of the invention, as indicated by the appended claims, is intended to include variations from the embodiments provided which are nevertheless described by the broad meaning and range properly afforded to the language of the claims, or to the equivalents thereof.

CLAIMS

I claim:

1. A process for treatment of an aqueous feed stream in heat transfer equipment, said heat transfer equipment comprising at least one evaporator with at least one heat transfer surface, to produce a low solute containing distillate stream and a high solute/solids containing blowdown stream, in which said feed stream has 5 minimal tendency to scale said heat transfer surface, said process comprising:

(a) providing a feed water stream containing soluble and insoluble inorganic and organic species therein, said species comprising:

(II) multi-valent metal cations,

(III) alkalinity,

10 (IV) at least one molecular species which is at low ionization levels when in solution at around neutral pH;

(b) removing a portion or substantially all multi-valent cations metal from said feed stream, and

15 (c) reducing the tendency of said feed water or form scale on said heat transfer surfaces, when said feed water is concentrated to a selected concentration factor at a selected pH, by effecting, in any order, one or more of the following:

(I) removing substantially all alkalinity from said feed water stream;

20 (II) removing dissolved gas from said feed water stream;

(III) raising the pH of said feed water stream to at least about 9 or higher;

(d) passing the product from step (c) into said heat transfer equipment, wherein said heat transfer equipment:

25 (I) contains a plurality of heat transfer surfaces,

(II) contains a circulating high solids solution, and

(III) wherein the pH of said circulating solution is maintained to at least about 9, or higher,

(e) so as to concentrate said feed water to said selected concentration factor, to produce:

- (I) a high solute/solids containing blowdown stream, and
- (II) a low solute containing distillate stream.

2. The process as set forth in claim 1, wherein the step of removing said multi-valent cations includes removing substantially all said alkalinity associated with hardness, and is accomplished in a single unit operation.

3. The process as set forth in claim 2, wherein said single unit operation comprises a weak acid cation ion exchange system operated in a hydrogen form.

4. The process as set forth in claim 1(b), wherein the step of said multi-valent cation removal is accomplished in a weak acid cation ion exchange system that is operated in a sodium form.

5. The process as set forth in claim 1, further comprising the step of adding acid before the step of removing said dissolved gas, to effect conversion of alkalinity to carbon dioxide.

6. The process as set forth in claim 3, wherein said feed water stream contains more multi-valent cations than alkalinity, and further comprising, before feeding said feed water to said weak acid cation exchange system, the step of adjusting the ratio of multi-valent cations to alkalinity by adding a base to said feed water, so as to raise the alkalinity of said feed water.

7. The process as set forth in claim 3, wherein said feed water stream contains more alkalinity than multi-valent cations, and further comprising, before feeding said feed water to said weak acid cation exchange system, the step of addition of acid to said feed water, so as to remove the excess alkalinity in said feed water.

8. The process as set forth in claim 1, wherein the step of multi-valent cation removal is accomplished by passing said feed water through a sodium form strong acid cation ion exchange system.

9. The process as set forth in claim 1, wherein in step (c), the pH is raised to between 10 and 11.

10. The process as set forth in claim 1, wherein in step (c), the pH is raised to between 9 and 10.

11. The process as set forth in claim 10, wherein said sparingly ionized species when in neutral or near neutral pH aqueous solution comprises a weak acid with a pKa₁ of about 7.5 or higher.

12. The process as set forth in claim 1, wherein in step (c), the pH is raised to between 11 and 12.

13. The process as set forth in claim 1, wherein in step (c), the pH is raised to between 12 and 13.

14. The process as set forth in claim 1, wherein in step (c), the pH is raised to greater than or at least about 13.

15. The process as set forth in claim 1, wherein said low ionized species when in neutral or near neutral pH aqueous feed stream comprises silica (SiO₂).

16. The process as set forth in claim 1, wherein said low ionized species when in neutral or near neutral pH aqueous feed stream comprises meta/ortho silicic acid (H₄ SiO₄).

17. The process as set forth in claim 1, wherein said low ionized species when in neutral or near neutral pH aqueous feed stream comprises an ionizable organic carbon species.

18. The process as set forth in claim 1, wherein said low ionized species when in neutral or near neutral pH aqueous feed stream comprises boron, or derivatives thereof.

19. The process according to claim 1, wherein the step of removal of multi-valent cations is accomplished by addition of an alkali to simultaneously raise pH while precipitating hardness from said feed water stream.

20. The process as set forth in claim 1, further comprising the step of adding caustic before the step of removing dissolved gas, to effect removal of gasses such as ammonia.

21. The process as set forth in claim 1, wherein said at least one molecular species which is at low ionization levels when in solution at around neutral pH comprises silica, and wherein said blowdown stream contains silica up to about 160 ppm.

22. The process as set forth in claim 21, wherein said silica is present in said blowdown stream at up to about 2500 ppm.

23. The process as set forth in claim 21, wherein said silica is present in said blowdown stream at up to about 5000 ppm.

24. The process as set forth in claim 21, wherein said silica is present in said blowdown stream at up to about 7500 ppm.

25. The process as set forth in claim 21, wherein said silica is present in said blowdown stream at up to about 10,000 ppm.

26. The process as set forth in claim 21, wherein said silica is present in said blowdown stream at up to about 20,000 ppm.

27. The process as set forth in claim 21, wherein said silica is present in said blowdown stream at up to about 50,000 ppm.

28. The process as set forth in claim 21, wherein said silica is present in said blowdown stream at up to about 75,000 ppm.

29. The process as set forth in claim 21, wherein said silica is present in said blowdown stream at up to about 100,000 ppm.

30. The process as set forth in claim 1, wherein said feed water stream comprises silica, and wherein said heat transfer equipment is operated without limitation of the concentration of silica present in said blowdown stream.

31. The process according to claim 1, wherein the ratio of the quantity of said distillate stream produced to the quantity of said feed water stream provided is greater than about 50%.

32. The process according to claim 1, wherein the ratio of the quantity of said distillate stream produced to the quantity of said feed water stream provided is greater than about 85%.

33. The process according to claim 1, wherein the ratio of the quantity of said distillate stream produced to the quantity of said feed water stream provided is greater than about 95%.

34. The process according to claim 1, wherein the ratio of the quantity of said distillate stream produced to the quantity of said feed water stream provided is greater than about 99%.

35. The process according to claim 1, wherein said heat transfer equipment comprises falling thin film evaporation equipment, operating as a single unit, or operating in series, or operating in parallel to generate said distillate stream and said blowdown stream.

36. The process according to claim 1, wherein said heat transfer equipment comprises forced circulation evaporation equipment operating as a single unit or operating in parallel to generate said distillate stream and a high solids blowdown stream.

37. The process according to claim 1, wherein said heat transfer equipment comprises natural circulation evaporation equipment operating as a single unit or operating in parallel to generate said distillate stream and a high solids blowdown stream.

38. The process according to claim 35 or claim 36 or claim 37, wherein said heat transfer surfaces are tubular.

39. The process according to claim 35 or claim 36 or claim 37, wherein said heat transfer surfaces are plates.

40. The process as set forth in claim 38 or claim 39, wherein said heat transfer surfaces are operated in a vertical position.

41. The process as set forth in claim 38, wherein said heat transfer surfaces are operated in a horizontal position.

42. The process as set forth in claim 38 wherein said heat transfer surfaces are designed for enhanced heat transfer.

43. The process as set forth in claim 38 wherein said circulating solution is heated on the interior of the tubes.

44. The process as set forth in claim 38 or claim 39, wherein said circulating solution is heated on the exterior of the tubes or plates.

45. The process as set forth in claim 39, wherein said plates are die pressed plates.

46. The process as set forth in claim 39, wherein said plates are made from flat sheets welded together and then formed into final shape by internal pressure.

47. The process as set forth in claim 39, wherein said plates are fabricated as welded assemblies.

48. The process as set forth in claim 39, wherein said plates are gasketed.

49. The process as set forth in claim 35 or claim 36 or claim 37, wherein said heat transfer equipment is operated in a steam driven multiple effect mode.

50. The process as set forth in claim 35 or claim 36 or claim 37, wherein said heat transfer equipment is operated in a mechanical vapor recompression mode.

51. The process as set forth in claim 35 or claim 36 or claim 37, wherein said heat transfer equipment is operated in a thermal compression mode.

52. The process as set forth in claim 35 or claim 36 or claim 37, wherein said heat transfer equipment is operated as a multiple stage flash evaporator.

53. The process as set forth in claim 35 or claim 36 or claim 37, further comprising the step of treating said high solute concentrate stream in a crystallizer operating as a single unit or operating in parallel to generate said distillate stream and said high solids blowdown stream.

54. The process as set forth in claim 36 or claim 53, comprising the step of further treating said high solids blowdown stream after reaching said selected concentration factor in a solids dewatering device operating as a single unit or operating in parallel.

55. The process as set forth in claim 54, further comprising the step of generation of a high solute low suspended solids filtrate water stream, and still further comprising the step of directing said low filtrate water stream to the inlet of said heat transfer equipment for further processing.

56. The process as set forth in claim 55, further comprising the step of generation of a dry solids product for disposal.

57. The process as set forth in claim 1, wherein the step of removing said dissolved gases, is further comprised of lowering the pH of said feed stream to remove any remaining alkalinity and release carbon dioxide.

58. The process as set forth in claim 57, further comprised of heating said acidified feed water stream to enhance gas removal in a degasifier prior to entering said heat transfer equipment.

59. The process as set forth in claim 1, wherein step (d) further comprising distributing said circulating solution across one side of said plurality of heat transfer surfaces to generate a steam vapor.

60. The process as set forth in claim 59, further comprising collecting said steam vapor and slightly compressing it to form a compressed steam vapor.

61. The process as set forth in claim 60, further comprising directing said compressed steam vapor to a second side of said plurality of heat transfer surfaces to condense said compressed steam vapor into said distillate stream.

62. The process according to claim 1, wherein the step of raising the pH is accomplished by addition of a base in aqueous solution, said base selected from the group consisting of (a) sodium hydroxide, (b) sodium carbonate, (c) potassium hydroxide, and (d) potassium carbonate.

63. The process according to claim 1, wherein the step of raising the pH is accomplished by addition of an aqueous organic base.

64. The process according to claim 1, wherein the ratio of the quantity of said distillate stream produced to the quantity of said feed water stream provided is between 90 and 98 percent.

65. The process according to claim 1, wherein said feed water stream further comprises cooling tower blowdown.

66. The process according to claim 1, wherein said feed water further comprises scrubber blowdown.

67. The process according to claim 1, wherein said feed water further comprises water utilized in ash transport in a coal fired steam-electric power plant.

68. The process according to claim 1, wherein said feed water stream comprises ash pond water.

69. The process according to claim 1, wherein said feed water stream comprises ash-slubbing water.

70. The process according to claim 1, wherein said feed water stream comprises effluent from sewage treatment.

71. The process according to claim 1, wherein said feed water stream comprises effluent from a food processing treatment.

72. The process according to claim 1, wherein said feed water stream comprises boiler blowdown.

73. The process according to claim 1, wherein said feed water stream comprises a concentrated stream from membrane separation equipment.

74. The process according to claim 1, wherein said feed water stream comprises effluent from oil refining operations.

75. The process as set forth in claim 35 or claim 36 or claim 37, comprising the step of further treating said high solids blowdown stream after reaching said selected concentration factor in a spray dryer to dry solids.

76. The process as set forth in claim 1, wherein the step of multi-valent cation removal is partially accomplished by passing said feed water stream through membrane softening equipment.

77. The process as set forth in claim 1, wherein the step of multi-valent cation removal is accomplished by increasing the pH to at least about 10 in said feed water stream and passing the pH adjusted stream through membrane separation equipment to filter out hardness precipitate.

78. In a process for the concentration of an aqueous feed stream in an evaporator to produce a low solute containing distillate stream, and retaining at least a portion of said aqueous feed stream in said evaporator to increase the concentration of a solute to a selected concentration factor at a selected pH in said retained portion 5 of said aqueous feed stream, the improvement which comprises feeding said evaporator with an aqueous feed stream characterized at the time of initial entry into said evaporator, by:

- (I) substantially no multi-valent cations,
- (II) substantially no alkalinity,
- 10 (III) substantially no dissolved or suspended gases, and
- (IV) a pH of at least 9 or above.

79. The process as set forth in claim 78 wherein greater than 80% of the multi-valent cations are removed from said feed stream.

80. The process as set forth in claim 78 wherein greater than 90% of the multi-valent cations are removed from said feed stream.

81. The process as set forth in claim 78 wherein greater than 98% of the multi-valent cations are removed from said feed stream.

82. The process as set forth in claim 78 wherein greater than 80% of the alkalinity is removed from said feed stream.

83. The process as set forth in claim 78 wherein greater than 90% of the alkalinity is removed from said feed stream.

84. The process as set forth in claim 78 wherein greater than 98% of the alkalinity is removed from said feed stream.

85. The process as set forth in claim 78 wherein greater than 80% of the gases are removed from said feed stream.

86. The process as set forth in claim 78 wherein greater than 90% of the gases are removed from said feed stream.

87. The process as set forth in claim 78 wherein greater than 98% of the gases are removed from said feed stream.

88. The process as set forth in claim 78 wherein said bases are selected from the group consisting of carbon dioxide, ammonia, oxygen, nitrogen and mixtures thereof.

89. The process as set forth in claim 78, wherein said aqueous feed stream is further characterized by minimizing or eliminating scale inhibitor solution or scale dispersant solution in said aqueous feed stream.

90. A method for treating a feed water stream, in a least one evaporator system and for simultaneously (a) reducing the scaling potential and (b) allowing an increased heat transfer rate and (c) allowing higher recovery and (d) minimizing or eliminating scale control methods in an existing heat transfer system, to produce a low solute containing distillate stream and a high solute/solids containing blowdown stream, said method comprising:

(a) providing a feed water stream containing soluble and insoluble species therein, said species comprising two or more of the following:

(I) multi-valent metal cations,

(II) alkalinity, and

(III) at least one molecular species which is at low ionization levels when in solution at around neutral pH;

(b) removing a portion or substantially all multi-valent cations from said feed stream, and

15 (c) reducing the tendency of said feed water to form scale when
said feed water is concentrated to a selected concentration factor at a selected
pH, by effecting, in any order, one or more of the following;

(I) removing substantially all alkalinity from said feed water stream;

(II) removing dissolved or suspended gases from said feed water stream, whether initially present or created during said multi-valent cation or said alkalinity removal step or said pH adjustment step;

(III) raising the pH of said feed water stream to at least about 9 or higher;

(d) passing the product from step (c) into heat transfer equipment, wherein said heat transfer equipment:

(I) contains a plurality of heat transfer surfaces,

(II) contains a circulating high solutes/solids solution, and

(III) the pH of said circulating solution is maintained to at least 9, or higher,

(e) so as to concentrate said feed water to said selected concentration factor, to produce:

(I) a high solute/solids containing blowdown stream, and

(II) a low solute containing distillate stream.

91. In a process for the purification of an aqueous feed stream comprising solutes and solvent by using evaporation equipment to increase the concentration of said aqueous stream to a selected concentration factor by generating a low solute containing distillate stream and retaining at least a portion of said aqueous feed stream in said evaporation equipment to increase the concentration of a selected solute to a selected concentration factor in said retained portion of said aqueous feed stream, the improvement which comprises controlling solutes, multi-valent metal cations, alkalinity, and carbon dioxide in said aqueous feed stream to a level where

the tendency to form scale is effectively eliminated at said selected concentration

10 factor, by

(a) prior to feeding of said aqueous feed stream to said evaporation equipment, in any order,

(I) minimizing multi-valent cations in said aqueous feed stream,

(II) minimizing alkalinity of said aqueous feed stream,

(III) minimizing gases dissolved or suspended in said aqueous feed stream;

(b) then, after step (a), increasing the pH of said aqueous feed stream in said evaporation equipment to at least about 9, or higher.

92. The process as set forth in claim 1, or claim 78, or claim 90, or claim 91, further comprising, during the step of removing alkalinity, the additional step of removing substantially all non-hydroxide alkalinity not associated with hardness.

93. The process as set forth in claim 90, or claim 91, wherein the step of raising the pH of said feed water stream, comprises raising the pH to between about 10 and about 11.

94. The process as set forth in claim 90, or claim 91, wherein the step of raising the pH of said feed water stream comprises raising the pH to between about 11 and about 12.

95. The process as set forth in claim 90, or claim 91, wherein the step of raising the pH of said feed water stream comprises raising the pH to between about 12 and about 13.

96. The process according to claim 1, wherein said feed water stream comprises effluents from hydrocarbon recovery operations as produced water.

97. The process as set forth in claim 90, or claim 91, or claim 91, wherein during the step of raising the pH of said feed water stream comprises raising the pH to between about 9 and about 10.

98. The process as set forth in claim 19, wherein a high solids containing waste stream is generated and, further comprising, de-watering of said high solids containing waste stream.

99. The process as set forth in claim 98, wherein a low suspended solids stream is generated and, further comprising directing said low suspended solids stream back to the inlet of said softener.

100. The process as set forth in claim 90, or claim 91, wherein the step of raising the pH of said feed water stream comprises raising the pH to greater than or at least about 13.

101. The process as set forth in claim 1, wherein the steps of (b) removing multi-valent cations, and (c) removing alkalinity, removing dissolved gases, and increasing pH are accomplished prior to a membrane process to pre-concentrate the feed stream upstream of said heat transfer equipment described under step (d).

102. The process as set forth in claim 1, wherein the removal of multi-valent cations and partially raising the pH are accomplished prior to pre-concentrating said feed stream in a membrane process prior to step (c).

103. Apparatus for treatment of a feed water stream, said feed water stream characterized by the presence of two or more of the following:

(I) multi-valent metal cations,

(II) alkalinity,

5 (III) at least one molecular species which is at low ionization levels

when in solution at around neutral pH,

10 to produce a low solute containing distillate stream and a high solute/solids containing blowdown stream, said apparatus comprising:

a) pretreatment equipment for effectively eliminating the tendency of said feed water to form scale on heat transfer surfaces when said feed water is concentrated to a desired concentration factor at a selected pH, comprising, in any order:

(I) at least one softener for removing a portion or

15

substantially all multi-valent cations from said feed stream,
and one or more of the following:

20

(II) at least one de-alkalizer for removing essentially all
alkalinity from said feed water stream,
(III) a degasifier for removing dissolved gases,
(IV) chemical addition apparatus for raising the pH of
said circulating solution in said heat transfer equipment to a
selected pH of at least about 9 by adding a selected base thereto,
to urge said at least one molecular species with low ionization
levels when in solution at about neutral pH toward increased
ionization;

25

(b) one or more evaporator units, said one or more evaporator units,
treating said feed water to produce a high solute/solids containing blowdown
stream and a low solute containing distillate stream, and to concentrate said
feed water to said selected concentration factor.

104. The apparatus as set forth in claim 103, further comprising,
downstream of one or more said evaporator units, to further process said low solute
containing distillate stream therefrom, a cation exchange unit.

105. The apparatus as set forth in claim 103, further comprising,
downstream of one or more said evaporator units, to further process said low solute
containing distillate stream therefrom, an anion exchange unit.

106. The apparatus as set forth in claim 103, further comprising,
downstream of one or more said evaporator units, to further process said low solute
containing distillate stream therefrom, at least one mixed bed ion exchange unit.

5

107. The apparatus as set forth in claim 104 or claim 105 or claim 106,
further comprising an ion exchange resin regenerator that generates an ion exchange
regenerant stream, and further comprising means for directing said ion exchange
regenerant stream to the inlet of said degasifier unit in order to treat said ion
exchange regenerant stream in said evaporator.

108. The apparatus as set forth in claim 103, further comprising, downstream of one or more said evaporator units, to further process the said low solute containing distillate stream therefrom, a continuous electrodeionization unit to produce (a) a substantially solute free water stream and (b) a solute containing waste stream.

109. The apparatus as set forth in claim 108 further including means for directing said solute containing waste stream to the inlet of said degasifier for further processing.

110. The apparatus as set forth in claim 105, further comprising de-oiling apparatus upstream of said multi-valent cation removal softener.

111. The apparatus as set forth in claim 103, further comprising filtration equipment downstream of said softener.

112. The apparatus as set forth in claim 103, further including means for directing backwash water to the inlet of said softener for further processing.

Figure 1

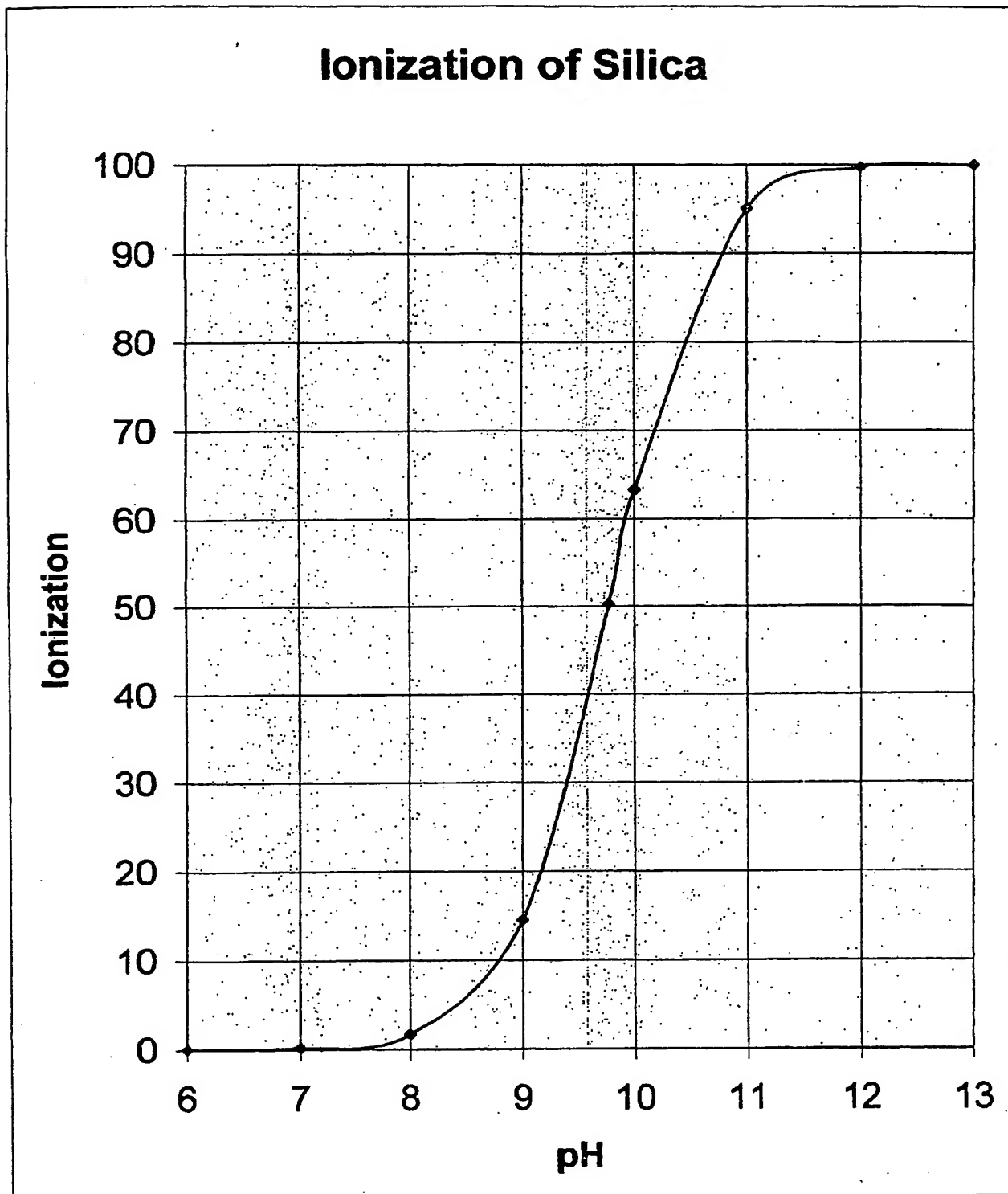


Figure 2

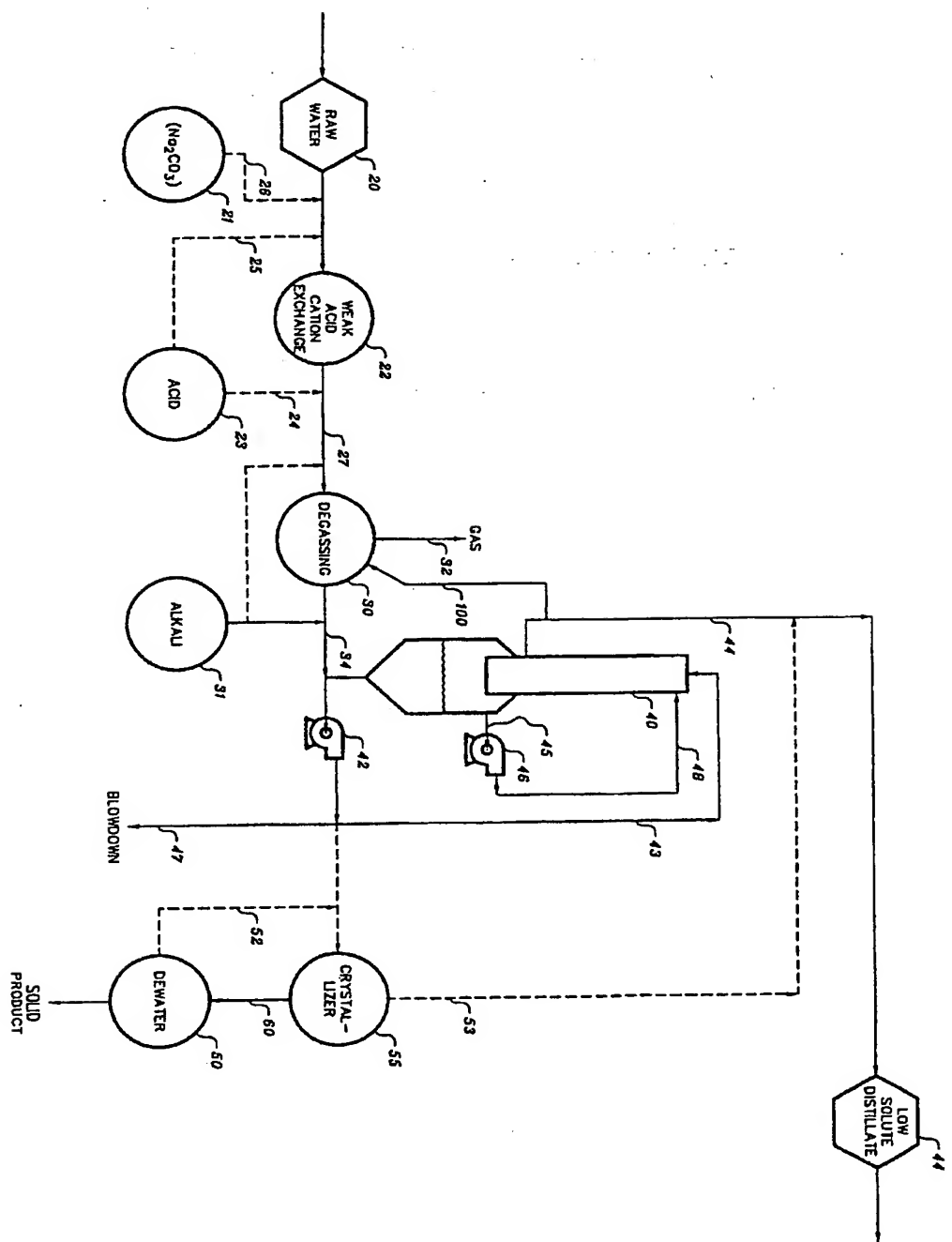


Figure 3

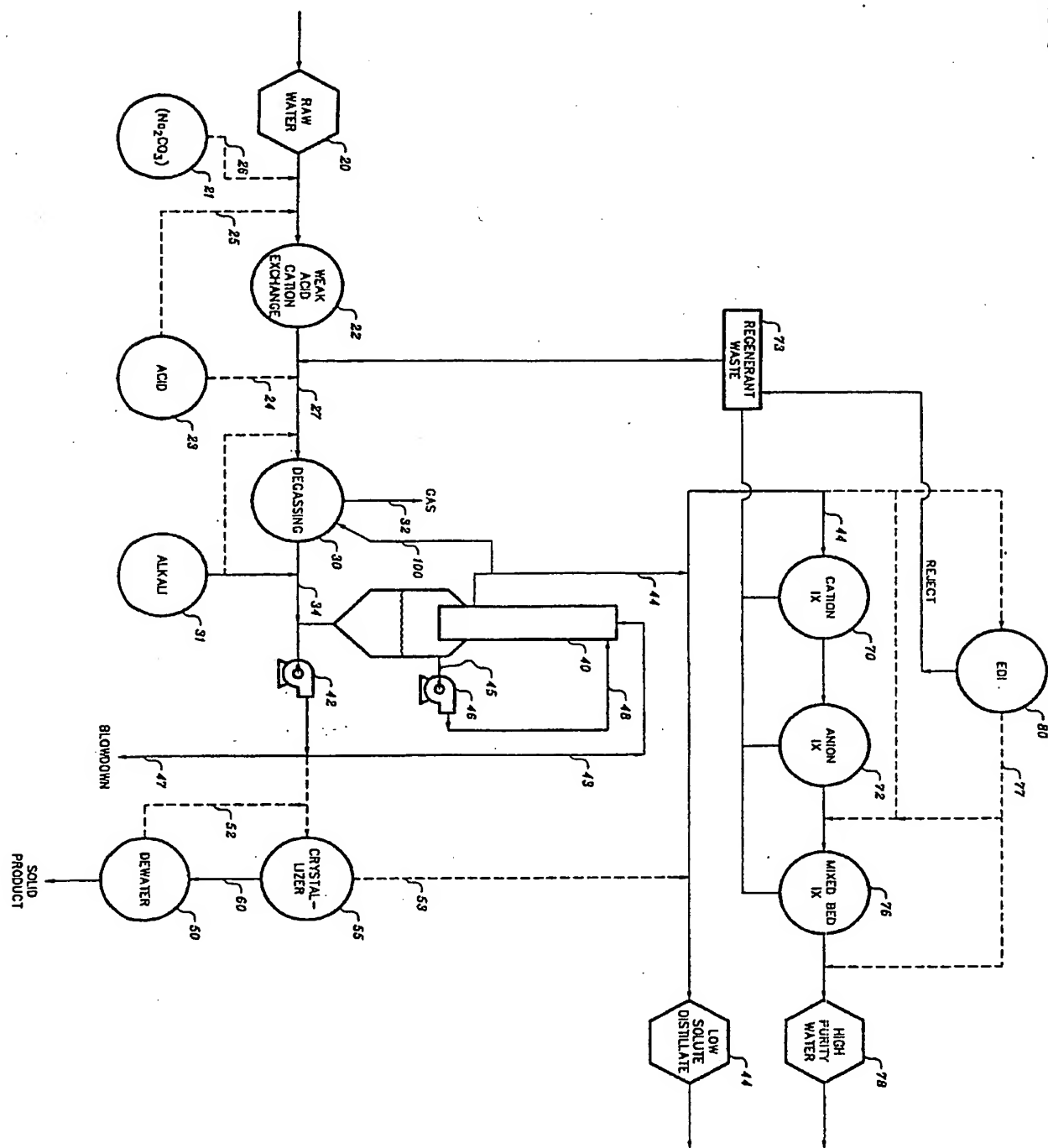


Figure 4

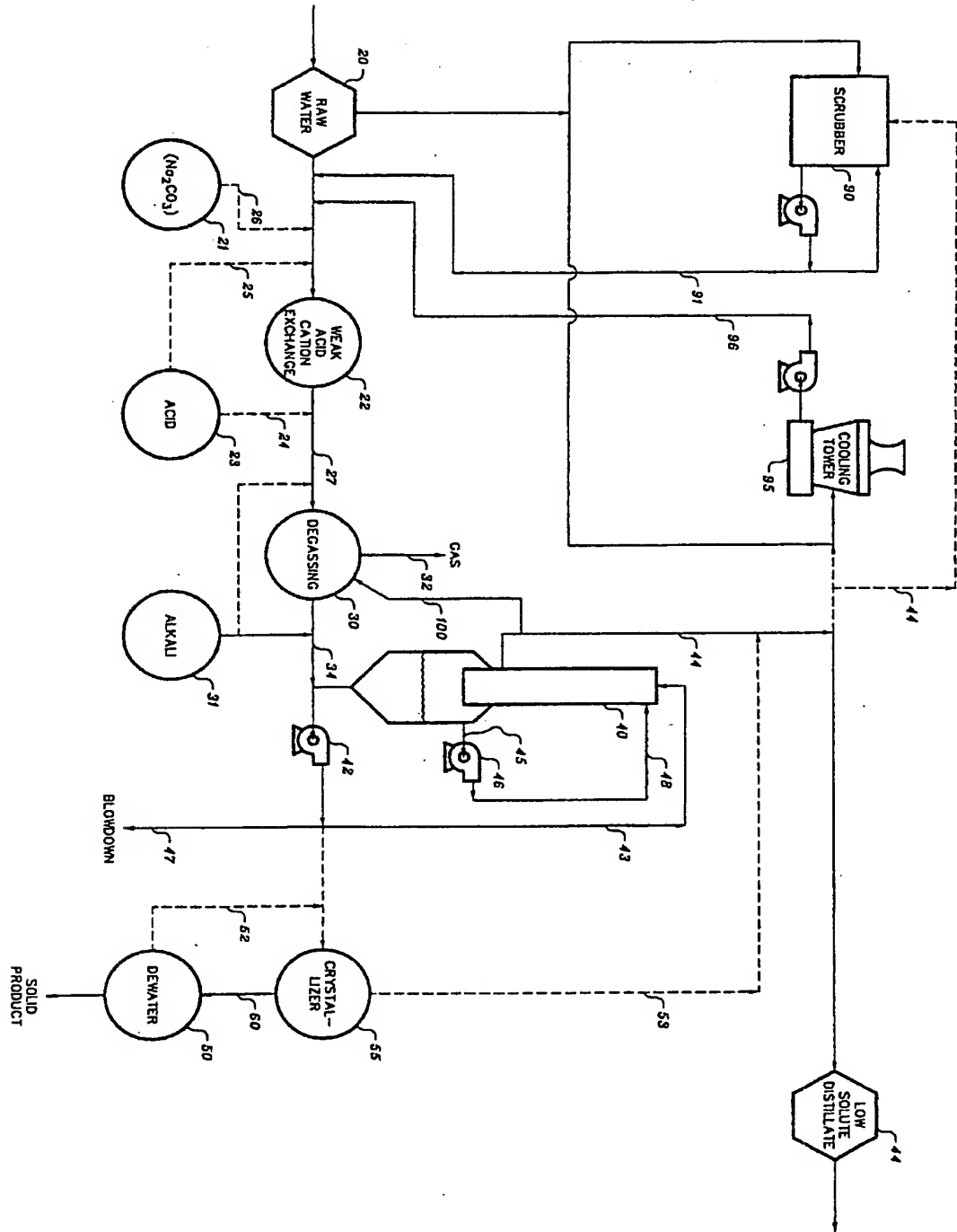


Figure 5

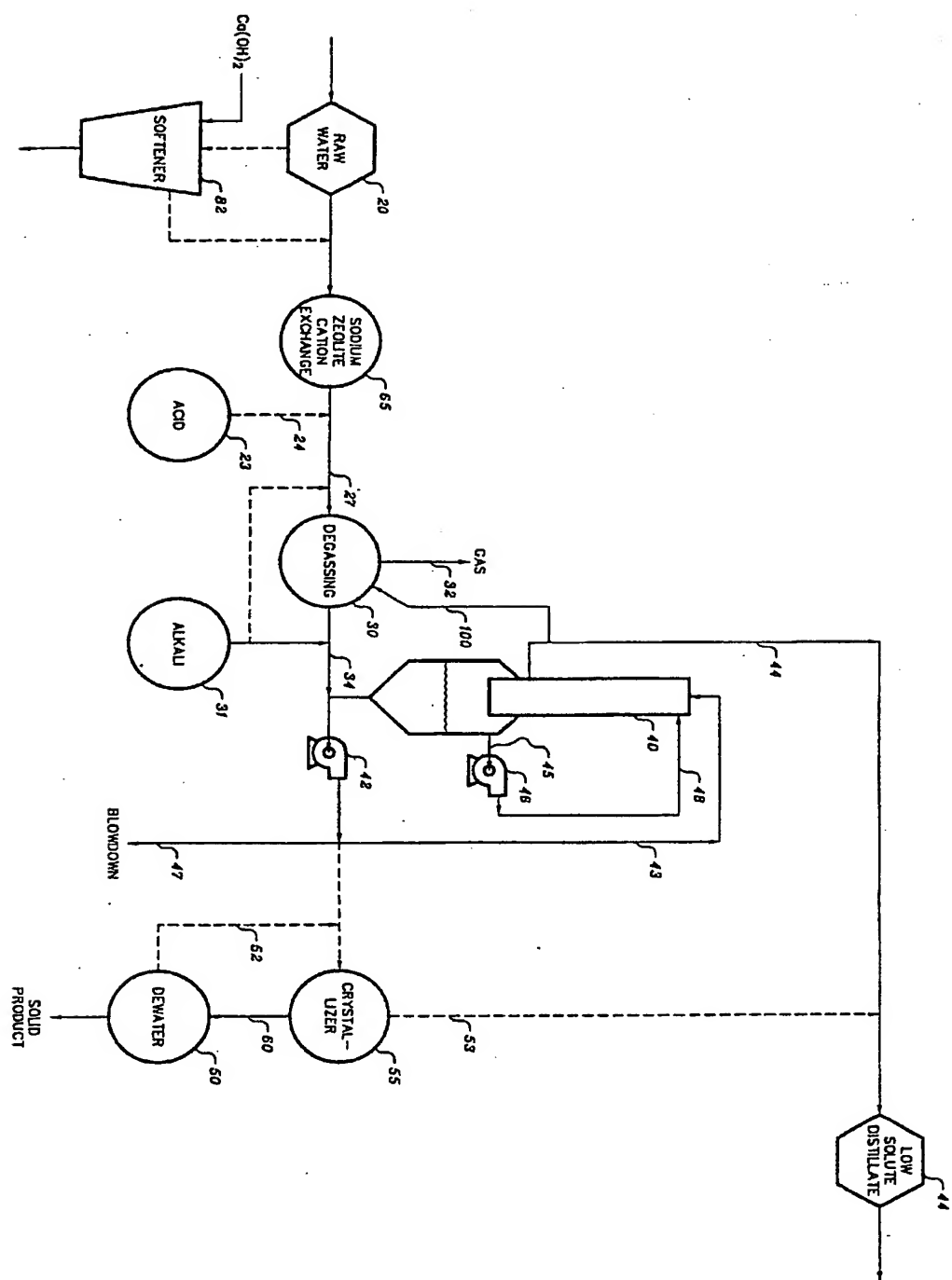
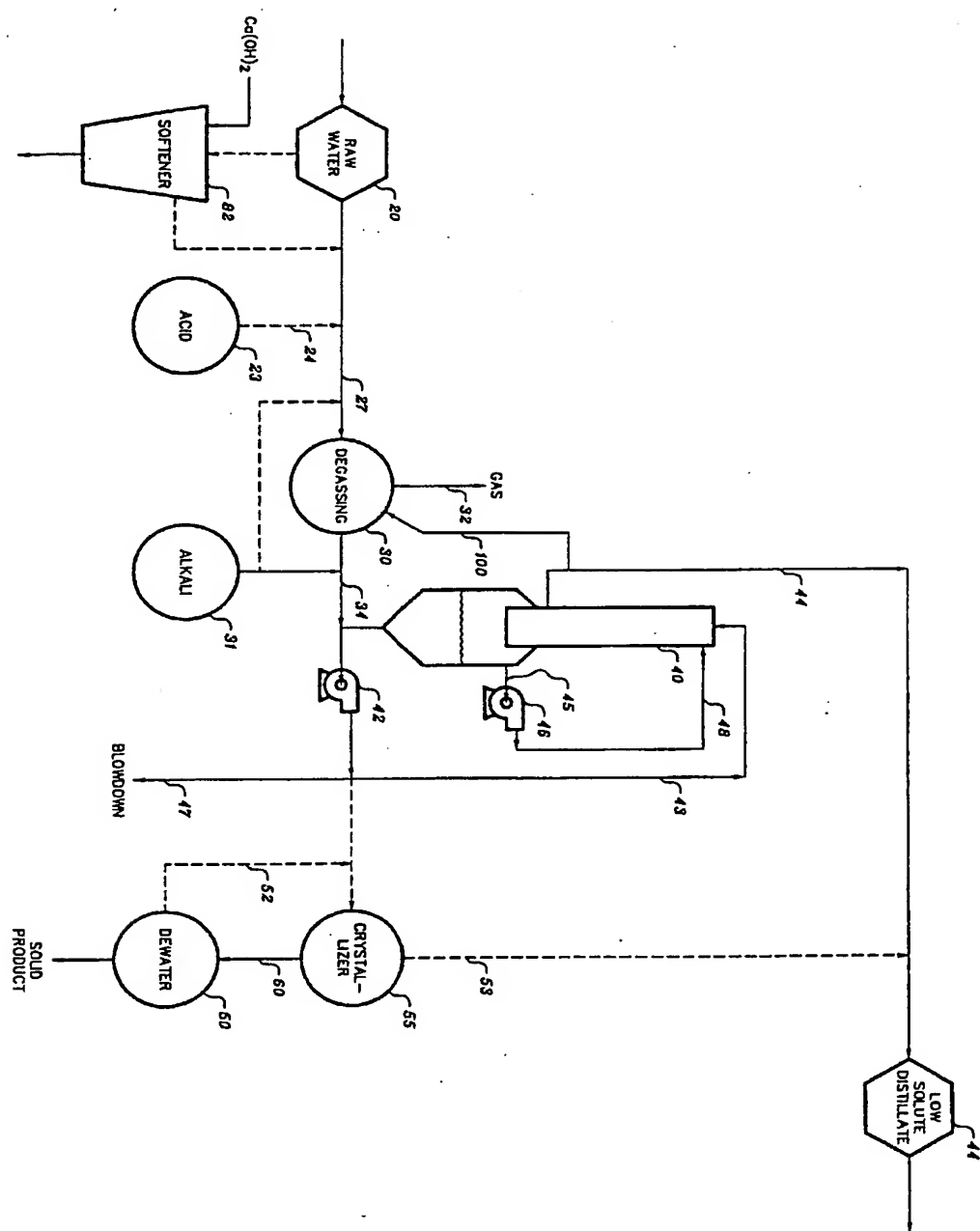


Figure 6



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PTO/SB/01 (8/96)	DECLARATION		Attorney Docket Number	4553-00013
	Declaration	OR	First Named Inventor	Keith R. Minnich
<input type="checkbox"/> Submitted with Initial Filing	<input checked="" type="checkbox"/> Submitted after Initial Filing		COMPLETE IF KNOWN	
			Application Number	
			Filing Date	
			Group Art Unit	
			Examiner Name	

As a below named inventor, I hereby declare that:

My residence, post office address, and citizenship are as stated below next to my name.

I believe I am the original, first and sole inventor (if only one name is listed below) or an original, first and joint inventor (if plural names are listed below) of the subject matter which is claimed and for which a patent is sought on the invention entitled:

Method and Apparatus for High Efficiency Evaporation Operation

(Title of the Invention)

the specification of which

is attached hereto

OR

was filed on (MM/DD/YYYY) 10/17/2003 as United States Application Number or PCT

International Number PCT/US03/033066 and was amended on (MM/DD/YYYY)

I hereby state that I have reviewed and understand the contents of the above identified specification, including the claims, as amended by any amendment specifically referred to above.

I acknowledge the duty to disclose information which is material to patentability as defined in 37 C.F.R. 1.56, including for continuation-in-part applications, material information which became available between the filing date of the prior application and the national or PCT international filing date of the continuation-in-part application.

I hereby claim foreign priority benefits under Title 35, United States Code §119(a)-(d) or §365(b) of any foreign application(s) for patent, inventor's or plant breeder's rights certificate(s), or §365(a) of any PCT international application which designated at least one country other than the United States of America, listed below and have also identified below, by checking the box, any foreign application for patent, inventor's or breeder's rights certificate(s), or of any PCT international application having a filing date before that of the application on which priority is claimed.

Prior Foreign Application Number(s)	Country	Foreign Filing Date (MM/DD/YYYY)	Priority Not Claimed	Copy Attached? YES	Copy Attached? NO
PCT/US2003/033066	WO	10/17/2003	<input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/>	<input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input checked="" type="checkbox"/>	<input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/>

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Application Number(s)	Filing Date (MM/DD/YYYY)	Additional provisional Application numbers are listed on a supplemental priority sheet attached hereto.

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Attorney Docket Number 4553-00013

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I hereby claim the benefit under Title 35, United States Code §120 of any United States application(s), or §365C of any PCT international application designated the United States of America, listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in the prior United States of PCT International application in the manner provided by the first paragraph of Title 35, United States Code §112. I acknowledge the duty to disclose information which is material to patentability as defined in Title 37, Code of Federal Regulations §1.56 which became available between the filing date of the prior application and the national or PCT international filing date of this application.

U.S. Parent Application Number	PCT Parent Number	Parent Filing Date (MM/DD/YYYY)	Parent Patent Number (if applicable)

Additional U.S. or PCT international application numbers are listed on a supplemental priority sheet attached hereto.

As a named inventor, I hereby appoint the following attorney(s) and/or agent(s) to prosecute this application and to transact all business in the Patent and Trademark Office connected therewith:

Customer Number: 26753

Name	Registration Number	Name	Registration Number
Daniel D. Fetterley	20,323	Joseph D. Kuborn	40,689
George H. Solveson	25,927	Jeffrey S. Sokol	35,686
Gary A. Essmann	29,376	Peter T. Holsen	54,180
Thomas M. Wozny	28,922	Aaron T. Olejniczak	54,853
Michael E. Taken	28,120	Christopher M. Scherer	50,655
Joseph J. Jochman, Jr.	25,058	William L. Falk	27,709

Additional attorney(s) and/or agent(s) named on a supplemental sheet attached hereto.

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	Suite 1100		
CITY	Milwaukee	STATE	Wisconsin
COUNTRY	U.S.A.	TELEPHONE	(414) 271-7590
		FAX	(414) 271-5770

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under §1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

Name of Sole or First Inventor: A petition has been filed for this unsigned inventor

Given Name (first and middle [if any]) Family Name or Surname

Keith R. Minnich

Inventor's Signature *Keith R. Minnich* Date *20 Apr 2006*

RESIDENCE: City Pewaukee State WI Country US Citizenship US

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Additional inventors are being named on supplemental sheet(s) attached hereto.

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Attorney Docket Number 4553-00013

DECLARATION				ADDITIONAL INVENTOR(S) Supplemental Sheet			
Name of Additional Joint Inventor, if any:				<input checked="" type="checkbox"/> A petition has been filed for this unsigned inventor Given Name (first and middle [if any]) Family Name or Surname			
Ramkumar				Karlupudi			
Inventor's Signature				Date			
RESIDENCE: City	Waukesha	State	WI	Country	US	Citizenship	US
POST OFFICE ADDRESS		1981 Foxcroft Lane					
City	Waukesha	State	WI	Zip	53189	Country	US
Name of Additional Joint Inventor, if any:				<input type="checkbox"/> A petition has been filed for this unsigned inventor Given Name (first and middle [if any]) Family Name or Surname			
Richard M.				Schoen			
Inventor's Signature <i>Richard M. Schoen</i>				Date <i>7/20/06</i>			
RESIDENCE: City	Hartland	State	WI	Country	US	Citizenship	US
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<input type="checkbox"/> Additional inventors are being named on supplemental sheet(s) attached hereto.							

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**DECLARATION
FOR UTILITY OR DESIGN
PATENT APPLICATION**

Declaration
 Submitted with
 Initial Filing

Declaration
 Submitted after
 Initial Filing

Attorney Docket Number	4553-00013
First Named Inventor	Keith R. Minnich
COMPLETE IF KNOWN	
Application Number	
Filing Date	
Group Art Unit	
Examiner Name	

As a below named inventor, I hereby declare that:

My residence, post office address, and citizenship are as stated below next to my name.

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Prior Foreign Application Number(s)	Country	Foreign Filing Date (MM/DD/YYYY)	Priority Not Claimed	Copy Attached?
				YES NO
PCT/US2003/033066	WO	10/17/2003	<input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/>	<input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input checked="" type="checkbox"/>

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Application Number(s)	Filing Date (MM/DD/YYYY)	Additional provisional application numbers are listed on a supplemental priority sheet attached hereto.

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Attorney Docket Number **4553-00013**

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U.S. Parent Application Number	PCT Parent Number	Parent Filing Date (MM/DD/YYYY)	Parent Patent Number (if applicable)

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Thomas M. Wozny	28,922	Aaron T. Olejniczak	54,853
Michael E. Taken	28,120	Christopher M. Scherer	50,655
Joseph J. Jochman, Jr.	25,058	William L. Falk	27,709

Additional attorney(s) and/or agent(s) named on a supplemental sheet attached hereto.

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I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under §1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

Name of Sole or First Inventor: A petition has been filed for this unsigned inventor

Given Name (first and middle [if any]) Family Name or Surname

Keith R. **Minnich**

Inventor's Signature Date

Residence: City State Country Citizenship
Pewaukee **WI** **US** **US**

Mailing Address
W291 N3821 Round Hill Circle

City State Zip Country
Pewaukee **WI** **53072** **US**

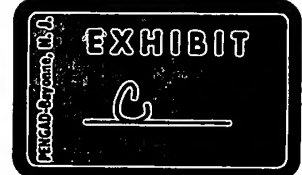
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Attorney Docket Number	4553-00013
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DECLARATION	ADDITIONAL INVENTOR(S) Supplemental Sheet
-------------	--

Name of Additional Joint Inventor, if any:		<input type="checkbox"/> A petition has been filed for this unsigned inventor	
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Inventor's Signature		Date	
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City Lake Mary	State FL	Zip 32746	Country US
Name of Additional Joint Inventor, if any:		<input type="checkbox"/> A petition has been filed for this unsigned inventor	
Given Name (first and middle [if any]) Richard M.		Family Name or Surname Schoen	
Inventor's Signature		Date	
Residence: City Hartland	State WI	Country US	Citizenship US
Mailing Address N67 W29767 Hartling Road			
City Hartland	State WI	Zip 53029	Country US
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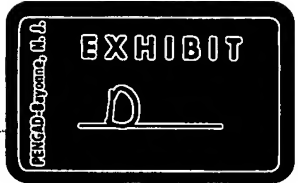
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Lisa Schollaert

From: Lisa Schollaert
Sent: Tuesday, March 13, 2007 11:00 AM
To: 'Ram Karlapudi'
Subject: RE: Assignment

Hi Ram

I am getting a little hassling from the attorney handling the patent
Please let me know if you will or will not be signing the declaration
I really hate to keep bothering you

Thanks so much

Regards

Lisa

-----Original Message-----

From: Lisa Schollaert
Sent: Wednesday, January 31, 2007 2:42 PM
To: 'Ram Karlapudi'
Subject: RE: Assignment

Hi Ram

As of today we have not received the signed declaration from you regarding our patent.
Could you please let me know when to expect it?

Thanks

Lisa

-----Original Message-----

From: Ram Karlapudi [mailto:Ram_Karlapudi@pall.com]
Sent: Thursday, January 04, 2007 2:44 PM
To: Lisa Schollaert
Cc: billf@andruslaw.com
Subject: RE: Assignment

Lisa,

My Home address:

1705 Cherry Ridge Drive
Lake Mary, FL 32746

Please note that I will be out of the country for the next couple of weeks.

Best Regards

Ramkumar Karlapudi

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Tracking number	799103916798	Reference	Lisa/RamKumar	Wrong Address?
Signed for by	K.KARLAPUDI	Destination	Lake Mary, FL	Reduce future mistal FedEx Address Check
Ship date	Mar 14, 2007	Delivered to	Residence	
Delivery date	Mar 15, 2007 12:48 PM	Service type	Standard Envelope	Tracking a FedEx Shipment?
		Weight	0.5 lbs.	Go to shipper login
Status	Delivered			

Date/Time	Activity	Location	Details
Mar 15, 2007	12:48 PM Delivered	Lake Mary, FL	
	10:27 AM On FedEx vehicle for delivery	LONGWOOD, FL	
	7:30 AM At local FedEx facility	LONGWOOD, FL	
	5:57 AM At dest sort facility	ORLANDO, FL	
	4:10 AM Departed FedEx location	MEMPHIS, TN	
	11:25 PM Arrived at FedEx location	MEMPHIS, TN	
Mar 14, 2007	8:23 PM Left origin	PITTSBURGH, PA	
	7:18 PM Picked up	PITTSBURGH, PA	
	3:10 PM Package data transmitted to FedEx		

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English



English



English



English

**Select format:** HTML Text Wireless**Add personal message:**

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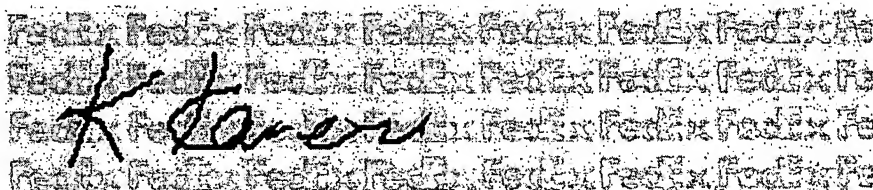
April 12, 2007

Dear Customer:

The following is the proof of delivery you requested with the tracking number 799103916798.

Delivery Information:

Status:	Delivered	Delivery location:	1705 CHERRY RIDGE DRIVE
			Lake Mary, FL 32746
Signed for by:	K.KARLAPUDI	Delivery date:	Mar 15, 2007 12:48
Service type:	Standard Envelope		



Shipping Information:

Tracking number:	799103916798	Ship date:	Mar 14, 2007
		Weight:	0.5 lbs.

Recipient:
Ramkumar Karlapudi
Residence
1705 Cherry Ridge Drive
Lake Mary, FL 32746 US

Reference

Shipper:
Beth Ann Wise
AQUATECH INTERNATIONAL CORP
1 FOUR COINS DRIVE
CANONSBURG, PA 15317 US
Lisa/RamKumar

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